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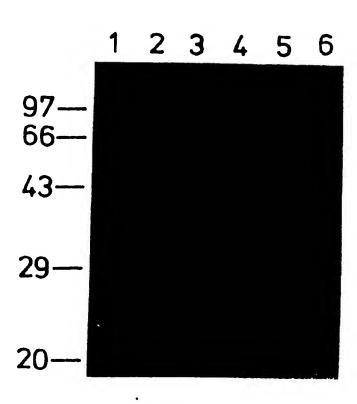
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(54) Title: PROTEIN PHOSPHATASE-1 CATALYTIC SUBUNIT INTERACTIONS

(57) Abstract

A method of identifying a compound which modulates the interaction between a PPIc and a regulatory subunit thereof, the method comprising determining whether a compound enhances or disrupts the interaction between (a) a PP1c or a fragment, variant, derivative or fusion thereof or a fusion of a fragment, variant or derivative and (b) a regulatory subunit which is able to bind to PP1c or a PP1c-binding fragment, variant, derivative or fusion of said subunit or a fusion of said fragment, variant or derivative. A method of affecting cellular metabolism or function, the method comprising administering to a cell (a) a compound which modulates the interaction between a PP1c and a regulatory subunit thereof or (b) a compound which mimics the effect of a regulatory subunit of PPIc or (c) a peptide capable of binding a PP1c and which affects the ability of PP1c to bind to a particular target and/or affects the regulation of PPIc activity, or a functional equivalent thereof.



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PROTEIN PHOSPHATASE-1 CATALYTIC SUBUNIT INTERACTIONS

The present invention relates to peptides and protein-protein interactions and to the use of peptides, peptide analogues and compounds which modulate proteinprotein interactions in the control of cellular metabolism and function.

Cellular metabolism or function is controlled by a number of regulatory agents, which are affected by extracellular factors, for example the physical condition of the cell or the binding of a messenger molecule to a receptor located on the cell surface. The extracellular factor may then initiate a cascade of secondary messenger reactions within the cell itself, leading ultimately to changes in some aspects(s) of metabolism or cell function.

It is well recognised by those skilled in the art that phosphorylation or dephosphorylation reactions often play a key role in regulating the activity of the proteins affected. Dephosphorylation reactions are catalysed by phosphatase enzymes, the activity of which may themselves be controlled by phosphorylation and/or dephosphorylation events. Whilst a substantial amount of knowledge has been accumulated regarding protein phosphatases as a group, the number and variety of these enzymes is such that detailed information concerning the mode of action of a specific phosphatase is not always available. There remains a need to further elucidate and characterise particular key enzymes.

The reversible phosphorylation of proteins regulates most aspects of cell life. About a third of all mammalian proteins are now thought to contain covalently bound phosphate and, since protein kinases and phosphatases probably account for approximately 2-3% of all human gene products (Hunter, 1995), many of these enzymes must typically phosphorylate/dephosphorylate numerous proteins in vivo. However, it is becoming increasingly clear that some protein kinases

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and phosphatases do not find their physiological substrates by simple diffusion within cells and that they are frequently directed to particular loci in the vicinity of their substrates by interaction with targeting subunits. In this way, the actions of protein kinases and phosphatases with inherently broad specificities are restricted and their properties tailored to the needs of a particular subcellular location, organelle or process (reviewed in Hubbard and Cohen, 1993; Faux and Scott, 1996).

Protein phosphatase-1 (PP1), one of the major protein serine/threonine phosphatases of eukaryotic cells, participates in the control of a variety of cellular functions that include glycogen metabolism, muscle contraction, the exit from mitosis (reviewed in [1,2]) and the splicing of mRNA [3]. However, evidence has been accumulating that different processes are regulated by distinct forms of PP1 in which the phosphatase catalytic subunit (PP1c) is complexed to specific "targeting subunits". These proteins not only direct PP1c to particular subcellular locations, but modify its specificity in unique ways and confer regulation by extracellular agonists (reviewed in [2,3]).

Several targeting subunits have been isolated and characterised, including the G_M -subunit that targets PP1c to both the glycogen particles and sarcoplasmic reticulum of striated muscles [4,5], the G_L subunit that targets PP1c to liver glycogen [6,7], the M-complexes responsible for the association of PP1c with the myofibrils of skeletal muscle [8,9] and smooth muscle [9-12], the p53 binding protein p53BP2 [13] and nuclear proteins such as sds22 [14] and NIPP1 [15,16]. PP1c is also reported to interact with other mammalian proteins such as the retinoblastoma gene product [17], ribosomal protein L5 [18], a 110 kDa nuclear protein that has yet to be identified [15] and two cytosolic proteins, termed inhibitor-1 and inhibitor-2. Inhibitor-1, and its homologue termed dopamine and cyclic AMP-regulated phosphoprotein (DARPP), become potent PP1 inhibitors after phosphorylation by cyclic AMP-dependent protein kinase

(PKA). Inhibitor-1 is thought to inactivate PP1c released from glycogen particles when G_M is phosphorylated by PKA [19]. Inhibitor-2 is present as a complex with PP1 in the cytosol, and there is evidence that one of its roles is to act like a molecular chaperone to ensure that the PP1 catalytic centre is folded correctly prior to its delivery to a specific targeting subunit [20]. It seems likely that many other PP1-targeting subunits will be identified over the next few years as a result of the introduction of powerful new techniques such as microcystin Sepharose affinity chromatography [8] and the yeast "two hybrid system" [13].

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The forms of PP1c isolated so far each contain a single PP1c-binding subunit, implying that the interaction of different targeting subunits with PP1c may be mutually exclusive. This, in turn, suggests that the binding sites for targeting subunits may overlap, and that the proportion of PP1 directed to any particular location may be determined by the amounts of each targeting subunit synthesised and their relative affinities for PP1. However, the different targeting subunits show surprisingly little similarity to one another. G_M and G_L are structurally related, yet display only 23% amino acid sequence identity over the first 286 residues of G_M , while G_L lacks the C-terminal 750 residues of G_M [7]. p53BP2 [13] and the M_{110} subunits from smooth muscle [10,11] and skeletal muscle [8] contain ankyrin repeats, but no other similarities have so far been detected between other PP1 targeting subunits.

The paradigm for the targeting subunit concept is protein phosphatase-1 (PP1), one of the major serine/threonine specific protein phosphatases of eukaryotic cells (Stralfors et al., 1985). This enzyme is involved in controlling diverse cellular functions including glycogen metabolism, muscle contraction, the exit from mitosis and the splicing of RNA (Cohen, 1989; Shenolikar, 1994; Wera and Hemmings, 1995). These different processes appear to be regulated by distinct PP1 holo-enzymes in which the same catalytic subunit (PP1c) is

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complexed to different targeting or regulatory subunits. The latter class of subunits act to confer *in vivo* substrate specificity not only by directing PP1c to the subcellular loci of its substrates, but also by enhancing or suppressing its activity towards different substrates. In addition, the regulatory subunits allow the activity of PP1 to be modulated by reversible protein phosphorylation and second messengers in response to extracellular stimuli.

Many regulatory subunits modulate the activity of PP1 towards its substrates. In the instance of the regulatory M_{110} subunit that targets PP1c to myosin, the region on the M_{110} subunit that enhances the dephosphorylation of myosin by PP1 has now been shown to be distinct from the region involved in targetting the PP1-M holoenzyme to myosin. These observations indicate that alterations in the substrate specificity of PP1c are likely to result from conformational changes induced by interactions with the targetting subunit and not simply as a direct result of targetting PP1c to its substrate. However, in the case of the glycogen binding subunit G_M , the dephosphorylation of glycogen phosphorylase and glycogen synthase was enhanced only under conditions when both the PP1- G_M complex and its substrates were bound to glycogen (Hubbard and Cohen, 1989) suggesting that targetting alone may be sufficient to enhance specificity.

Whilst the identity of the PP1-binding site(s) on any targeting subunit is unknown, it has now been realised that the control of the substrate specificity and activity of this key regulatory enzyme and its interactions are of therapeutic importance. Disruption of PP1-targeting subunit interactions provide a way of altering selectively the state of phosphorylation, and hence the activities, of particular PP1 substrates. We have now identified relatively small peptides from the G_M and M_{110} -subunits that interact with PP1, and which either disrupt or mimic the distinctive properties of myofibrillar and glycogen-associated forms of PP1. The binding of the G-subunit and the M-subunit of PP1 has also

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been found to be mutually exclusive.

A first aspect of the invention provides a method of identifying a compound which modulates the interaction between a PP1c and a regulatory subunit thereof, the method comprising determining whether a compound enhances or disrupts the interaction between (a) a PP1c or a fragment, variant, derivative or fusion thereof or a fusion of a fragment, variant or derivative and (b) a regulatory subunit which is able to bind to PP1c or a PP1c-binding fragment, variant, derivative or fusion of said subunit or a fusion of said fragment, variant or derivative.

Conveniently, the PP1c or a fragment, variant or derivative or fusion thereof or a fusion of a fragment, variant or derivative is one that is produced using recombinant DNA technology. By "fragment, variant, derivative or fusion of PP1c" we mean any such fragment, variant, derivative or fusion that retains the ability to interact with a regulatory subunit or a suitable PP1c-binding fragment, variant, derivative or fusion of said subunit or a fusion of said fragment, variant or derivative.

- By "regulatory subunit" we mean any such regulatory subunit. Further subunits are being identified all of the time. It is preferred if the regulatory subunit contains the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe as described below.
- By "PP1c-binding fragment, variant, derivative or fusion of said subunit or a fusion of said fragment, variant or derivative" we include any such fragments, variants, derivatives and fusions which are able to bind to PP1c. Conveniently, the fragments, variants, derivatives are made using recombinant DNA technology or, in the case of peptides and peptide derivatives and analogues they may be made using peptide synthetic methods.

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The enhancement or disruption of the interaction between the said PP1c or a fragment, variant, derivative or fusion thereof or a fusion of a fragment, variant or derivative and the said regulatory subunit or a fragment, variant, derivative or fusion thereof or a fusion of a fragment, variant or derivative can be measured *in vitro* using methods well known in the art of biochemistry and including any methods which can be used to assess protein-protein, protein-peptide and protein-ligand interactions.

The said interaction can also be measured within a cell, for example using the yeast two-hybrid system as is well known in the art.

It should be appreciated that before the present invention the dissociation of a PP1c-regulatory subunit has not been achieved using a small molecule such as a peptide or a peptide analogue or derivative. Thus, it is preferred if the compounds screened in the method of the first aspect of the invention are small molecules and in particular that they are not intact regulatory subunits of PP1c.

By "small molecule" we include any compounds which have a molecular weight of less than 5000, preferably less than 2000 and more preferably less than 1000. Conveniently, the compounds screened are compounds which are able to enter a cell either passively *via* the cell membrane or *via* an active uptake system.

A second aspect of the invention provides a method of identifying a compound which mimics the effect of a regulatory subunit of PP1c, the method comprising contacting said compound with PP1c and determining whether, in the presence of the compound, PP1c adopts the function of properties of a PP1c in the presence of a given regulatory subunit.

30 By "mimics the effect of a regulatory subunit of PP1c" we include the meaning

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that the compound modifies a property of PPlc in such a way that PP1c acts, in at least one respect, like PP1c that is interacting with a regulatory subunit.

Examples of the properties of PP1c that may be modified, and examples of compounds which modify the properties of PP1c which are therefore identifiable in this method are given below.

Preferably, in the methods of the first and second aspects the said regulatory subunit of PP1c is any one of M_{110} , G_L , G_M , M-complexes, p53 BP2, sds22, NIPPI, L5, Inhibitor-1, Inhibitor-2, or DARPP.

More preferably, the regulatory subunit of PP1c is any one of M_{110} , G_L , G_M , M-complexes or p53BP2, and still more preferably the regulatory subunit of PP1c is M_{110} or G_M .

In relation to the method of the first aspect of the invention the fragment of a regulatory subunit which is able to bind to PP1c is any of the peptides [G63-T93], [G63-N75], [E2-R575], [E2-P243], [E2-D118], [H100-P350] and peptide 63-80 of G_M or functional equivalents thereof or peptides comprising said peptide sequences provided that they are not the complete G_M regulatory subunit. Preferably the peptides are not [E2-R575] or [H100-P350].

As is described in more detail in the Examples, these peptides have been shown to bind to PP1c and it is convenient, in some circumstances, for the method to be carried out such that one of these peptide is displaced from, or the binding is enhanced to, PP1c. Suitably, the peptide may be labelled in a detectable manner to facilitate the detection of the interaction with PP1c. Conveniently, the peptide is labelled radioactively or fluorescently using methods well known in the art.

Also in relation to the method of the first aspect of the invention the fragment of a regulatory subunit which is able to bind to PP1c is any of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M_{110} or functional equivalents thereof or peptides comprising said peptide sequences provided that they are not the complete M_{110} regulatory subunit.

As is shown in more detail in the Examples these peptides have been shown to bind to PP1c.

- Also in relation to the first aspect of the invention the PP1c-binding fragment, variant or derivative of said regulatory subunit or a fusion of said fragment, variant or derivative comprises the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any amino acid.
- We have found that, surprisingly, many regulatory subunits that bind to PP1c contain the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any amino acid, preferably a naturally occurring amino acid. Typically, the PP1c-binding fragment, variant or derivative of said regulatory subunit or a fusion of said fragment, variant or derivative is a peptide (typically 8-400 amino acid residues, preferably 8-200, more preferably 8-10 and still more preferably 8-20 amino acid residues in length which comprises the given consensus peptide sequence).
- It is preferred if the PP1c-binding fragment, variant or derivative comprises, in addition to the said consensus peptide sequence, at least one basic residue in the four residues N-terminal of the consensus peptide sequence. Preferably, there are at least two basic residues in this position, more preferably at least three such residues.
- 30 It is also preferred wherein in the consensus peptide sequence Xaa is not Asp

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or Glu because the negative charge is believed to interfere with binding to PP1c. Similarly, it is preferred if Xaa is not a large hydrophobic residue such as Phe, Tyr, Trp, Ile or Leu.

- It is particularly preferred if the PP1c-binding fragment is a fragment of a regulatory subunit comprising the said consensus peptide sequence and therefore the peptide sequences which flank the consensus peptide sequence are the same as in the native regulatory subunit.
- Preferably the PP1c-binding fragment is a fragment of any of the M₁₁₀, G_L, G_M, M-complexes, p53BP2, sds22, NIPPI, L5, Inhibitor-1, Inhibitor-2 or DARPP regulatory subunits comprising said consensus sequence.

Although the methods of the first and second aspects of the invention do not rely on any particular mechanism whereby the modulation or mimicking occurs, it is preferred if the compound binds to a PP1c. Alternatively, but still preferably, the compound binds to a regulatory subunit of PP1c.

A further aspect of the invention provides a compound identifiable by the method of the first or second aspects of the invention.

A further aspect of the invention provides a compound which modulates the interaction between a PP1c and a regulatory subunit thereof said compound comprising any of the peptides [G63-T93], [G63-N75], [E2-R575], [E2-P243], [E2-D118], [H100-P350] and peptide 63 to 80 of G_M or functional equivalents or said compound comprising any of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M₁₁₀ or functional equivalents thereof or said compound comprising the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any naturally occurring amino acid or functional equivalents thereof, provided that the said compound is not a complete regulatory subunit

H100-P350].

ates the interaction ally the same way.

which conservative tution is intended Asn, Gln; Ser, Thr; to be a peptide with zely to enter a cell.

hich are resistant to are blocked, or both more of the peptide stantially the same is more resistant to

peptides [G63-T93], 350] or peptide 63 to bound consists of the -Y496] of M₁₁₀ or is not [E2-R575] or

hod of identifying a Plc and a regulatory egulatory subunit, the

method comprising selecting a compound which is capable of adopting the same or substantially the same conformation as a peptide bound to the regulatory subunit-binding site of PP1c or the same or substantially the same conformation as the portion of PP1c which binds to said peptide. Suitably, the peptide comprises the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any amino acid, preferably a naturally occurring amino acid. Conveniently, the said peptide consists of residues 63 to 75 of G_M.

It is particularly preferred if the conformation of the said peptide and the conformation of the said portion of PP1c is as defined by reference to the atomic coordinates given in Table A (see also Example 2). Example 2 provides further details of the peptide - PP1c interactions.

Table A provides the atomic coordinates for the given PP1c-peptide crystal structure.

A further aspect of the invention provides a compound identifiable by the aforementioned method of the invention.

It will be appreciated that the aforementioned compounds and peptides will be useful in medicine and, accordingly, the invention includes pharmaceutical compositions of the said compounds in combination with a pharmaceutically acceptable carrier.

The formulations may conveniently be presented in unit dosage form and may be prepared by any of the methods well known in the art of pharmacy. Such methods include the step of bringing into association the active ingredient (compound of the invention) with the carrier which constitutes one or more accessory ingredients. In general the formulations are prepared by uniformly and intimately bringing into association the active ingredient with liquid carriers

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or finely divided solid carriers or both, and then, if necessary, shaping the product.

Formulations in accordance with the present invention suitable for oral administration may be presented as discrete units such as capsules, cachets or tablets, each containing a predetermined amount of the active ingredient; as a powder or granules; as a solution or a suspension in an aqueous liquid or a non-aqueous liquid; or as an oil-in-water liquid emulsion or a water-in-oil liquid emulsion. The active ingredient may also be presented as a bolus, electuary or paste.

A tablet may be made by compression or moulding, optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing in a suitable machine the active ingredient in a free-flowing form such as a powder or granules, optionally mixed with a binder (eg povidone, gelatin, hydroxypropylmethyl cellulose), lubricant, inert diluent, preservative, disintegrant (eg sodium starch glycolate, cross-linked povidone, cross-linked sodium carboxymethyl cellulose), surface-active or dispersing agent. Moulded tablets may be made by moulding in a suitable machine a mixture of the powdered compound moistened with an inert liquid diluent. The tablets may optionally be coated or scored and may be formulated so as to provide slow or controlled release of the active ingredient therein using, for example, hydroxypropylmethylcellulose in varying proportions to provide desired release profile.

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Formulations suitable for topical administration in the mouth include lozenges comprising the active ingredient in a flavoured basis, usually sucrose and acacia or tragacanth; pastilles comprising the active ingredient in an inert basis such as gelatin and glycerin, or sucrose and acacia; and mouth-washes comprising the active ingredient in a suitable liquid carrier

Formulations suitable for parenteral administration include aqueous and non-aqueous sterile injection solutions which may contain anti-oxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the intended recipient; and aqueous and non-aqueous sterile suspensions which may include suspending agents and thickening agents. The formulations may be presented in unit-dose or multi-dose containers, for example sealed ampoules and vials, and may be stored in a freeze-dried (lyophilised) condition requiring only the addition of the sterile liquid carrier, for example water for injections, immediately prior to use. Extemporaneous injection solutions and suspensions may be prepared from sterile powders, granules and tablets of the kind previously described.

Preferred unit dosage formulations are those containing a daily dose or unit, daily sub-dose or an appropriate fraction thereof, of an active ingredient.

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It should be understood that in addition to the ingredients particularly mentioned above the formulations of this invention may include other agents conventional in the art having regard to the type of formulation in question, for example those suitable for oral administration may include flavouring agents.

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A further aspect of the invention provides a method of affecting cellular metabolism or function, the method comprising administering to a cell (a) a compound which modulates the interaction between a PP1c and a regulatory subunit thereof or (b) a compound which mimics the effect of a regulatory subunit of PP1c or (c) a peptide capable of binding a PP1c and which affects the ability of PP1c to bind to a particular target and/or affects the regulation of PP1c activity, or a functional equivalent thereof.

It will be appreciated that the said compounds are disclosed above with respect to specific compounds or with respect to methods of obtaining such compounds.

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In particular, it is preferred if the compound administered to the cell is any one or more of the peptides [G63-T93], [G63-N75], [E2-R575], [E2-P243], [E2-D118], [H100-P350] and peptide 63-80 of G_M or functional equivalents thereof or peptides comprising said peptide sequences or any one or more of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M_{110} or functional equivalents thereof or peptides comprising said peptide sequences. Preferably, the peptide is not [E2-R575] or [H100-P350].

In this embodiment it will be appreciated that functional equivalents include those compounds defined above as being functional equivalents, in particular, derivatives of peptides which are more readily able to enter a cell.

The compound may be administered to the cell in any suitable way, in particular in such a way that the compound will enter the cell in a suitable form to have its desired effect. Method of facilitating the entry of a compound into the cell are known in the art, for example, in relation to peptides the importins and penetrations may be used, or the peptides may be micro-injected or they may enter the cell in a suitable vehicle such as in a liposome.

20 Preferably, the cell is a cell in a mammalian body.

The aforementioned compounds of the invention or a formulation thereof may be administered by any conventional method including oral and parenteral (eg subcutaneous or intramuscular) injection. The treatment may consist of a single dose or a plurality of doses over a period of time.

Whilst it is possible for a compound of the invention to be administered alone, it is preferable to present it as a pharmaceutical formulation, together with one or more acceptable carriers. The carrier(s) must be "acceptable" in the sense of being compatible with the compound of the invention and not deleterious to

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the recipients thereof. Typically, the carriers will be water or saline which will be sterile and pyrogen free.

A further aspect of the invention provides a method of treating a patient in need of modulation of PP1c activity or function the method comprising administering to the patient an effective amount of a compound which modulates the interaction between a PP1c and a regulatory subunit thereof or (b) a compound which mimics the effect of a regulatory subunit of PP1c or (c) a peptide capable of binding a PP1c and which affects the ability of PP1c to bind to a particular target and/or affects the regulation of PP1c activity, or a functional equivalent thereof.

As will be apparent from what is described herein, protein phosphatase-1 (PP1) is one of the principal serine/threonine-specific protein phosphatases in human cells where it plays key roles in regulating a variety of physiological roles, including the metabolism of glycogen, the splicing of mRNA, the exit from mitosis and the contraction of smooth muscle. The different functions of PP1 are carried out by distinct species of this enzyme in which the same catalytic unit is complexed to different "targeting" subunits. The latter class of proteins direct PP1 to specific subcellular loci, tailor its properties to the needs of a particular locus and confer the ability to be regulated by extracellular signals (hormones, growth factors, neurotransmitters). Compounds as herein described that disrupt specific PP1-"targeting" subunits interactions or mimic the effect of a targeting subunit are likely to have a number of therapeutic uses as outlined below.

PP1 interacts with the M110-subunit which targets it to myosin in smooth muscle and enhances the rate at which PP1 dephosphorylates myosin. The dephosphorylation of myosin underlies the relaxation of smooth muscle. Thus compounds such as those disclosed herein which disrupt the interaction of PP1

with M110 in arterial muscle are expected to increase the phosphorylation of arterial myosin and elevate blood pressure.

The interaction of PP1 with M110 enhances the rate at which PP1 dephosphorylates myosin, but suppresses the rate at which it dephosphorylates glycogen phosphorylase. The disruption of the PP1-M110 interaction is therefore measured in a screen by looking for compounds which enhance the dephosphorylation of phosphorylase and/or suppress the dephosphorylation of the myosin P-light chain (see the Examples).

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Compounds, such as those disclosed herein, that mimic the effect of the M110 subunit in stimulating myosin dephosphorylation are expected to be useful in lowering blood pressure. Such compounds are identified by their ability to stimulate the dephosphorylation of the myosin P-light chain by the catalytic subunit of PP1. An example of such an assay, which shows that the N-terminal 38 residues of the M110 subunit stimulate the dephosphorylation of the myosin P-light chain by PP1, is shown in the Examples.

The interaction of PP1 with G_L targets the phosphatase to liver glycogen. This interaction enhances the dephosphorylation glycogen synthase which stimulates the conversion of glucose to glycogen. A compounds, such as those disclosed herein, disrupts the interaction between PP1 and G_L is expected to be useful in treating hypoglycaemia. The interaction of G_L with PP1 strongly suppresses the rate at which PP1 dephosphorylates glycogen phosphorylase. A compound, such as those disclosed herein, which disrupts the interaction of PP1 with G_L can be screened for very simply by its ability to increase the phosphorylase phosphatase activity of PP1 G_L . This can be carried out, for example, using rat liver glycogen pellet as described in the Examples. There is no need to use the purified enzyme.

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PP1 interacts with p53 BP2 (Helps et al, 1995) a protein which is known to interact with the tumour suppressor p53. The phosphorylation of p53 is known to enhance its ability to bind to DNA and hence its ability to function as a tumour suppressor. p53BP2 may be a protein which targets PP1 to p53 stimulating the dephosphorylation and inactivation of p53. A compound, such as those disclosed herein, which disrupts the interaction of PP1 with p53BP2 may enhance the phosphorylation of p53 and its ability to function as a tumour suppressor. Since p53BP2 suppresses the dephosphorylation of glycogen phosphorylase (Helps et al, 1995), compounds that disrupt the p53BP2-PP1 complex can be screened by measuring the increase in rate of dephosphorylation of glycogen phosphorylase.

The present invention provides peptides able to bind to the catalytic sub-unit of protein phosphatase-1 (hereinafter referred to as PP1c). Generally the peptides affect the ability of PP1c to bind to particular target(s) and/or the regulation of PP1c activity.

Peptides can be designed based on the sequences of regulatory subunits, especially in relation to the peptide consensus sequence found therein and its flanking sequences. Peptides can be synthesised by methods well known in the art. For example, peptides may be synthesised by the Fmoc-polyamide mode of solid-phase peptide synthesis as disclosed by Lu et al (1981) J. Org. Chem. 46, 3433 and references therein. Temporary N-amino group protection is afforded by the 9-fluorenylmethyloxycarbonyl (Fmoc) group. Repetitive cleavage of this highly base-labile protecting group is effected using 20% piperidine in N,N-dimethylformamide. Side-chain functionalities may be protected as their butyl ethers (in the case of serine threonine and tyrosine), butyl esters (in the case of glutamic acid and aspartic acid), butyloxycarbonyl derivative (in the case of cysteine) and 4-methoxy-2,3,6-trimethylbenzenesulphonyl derivative (in the case

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of arginine). Where glutamine or asparagine are C-terminal residues, use is made of the 4,4'-dimethoxybenzhydryl group for protection of the side chain amido functionalities. The solid-phase support is based on a polydimethylacrylamide polymer constituted from the three monomers dimethylacrylamide bisacryloylethylene diamine (cross (backbone-monomer), acryloylsarcosine methyl ester (functionalising agent). The peptide-to-resin cleavable linked agent used is the acid-labile 4-hydroxymethyl-phenoxyacetic acid derivative. All amino acid derivatives are added as their preformed symmetrical anhydride derivatives with the exception of asparagine and glutamine, which are added using a reversed N,N-dicyclohexyl-carbodiimide/1hydroxybenzotriazole mediated coupling procedure. All coupling and deprotection reactions are monitored using ninhydrin, trinitrobenzene sulphonic acid or isotin test procedures. Upon completion of synthesis, peptides are cleaved from the resin support with concomitant removal of side-chain protecting groups by treatment with 95% trifluoroacetic acid containing a 50% scavenger mix. Scavengers commonly used are ethanedithiol, phenol, anisole and water, the exact choice depending on the constituent amino acids of the peptide being synthesised. Trifluoroacetic acid is removed by evaporation in vacuo, with subsequent trituration with diethyl ether affording the crude peptide. Any scavengers present are removed by a simple extraction procedure which on lyophilisation of the aqueous phase affords the crude peptide free of Reagents for peptide synthesis are generally available from scavengers. Calbiochem-Novabiochem (UK) Ltd, Nottingham NG7 2QJ, UK. Purification may be effected by any one, or a combination of, techniques such as size exclusion chromatography, ion-exchange chromatography and (principally) reverse-phase high performance liquid chromatography. Analysis of peptides may be carried out using thin layer chromatography, reverse-phase high performance liquid chromatography, amino-acid analysis after acid hydrolysis and by fast atom bombardment (FAB) mass spectrometric analysis.

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The peptides may be derived from the targeting subunit(s) of PP1c, in particular from the subunits G_L , G_M , M_{110} and/or M_{21} . Additionally the peptides may be derived from other subunits such as different M-complexes, p53BP2, sds22, NIPP1, L5, Inhibitor-1, Inhibitor-2, DARPP or the like. Functional equivalents or portions of these peptides may also be used.

In a further aspect the present invention provides the use of peptides derived from targeting subunit(s) of PP1c, functional equivalents or portions thereof to affect cellular metabolism or function.

In a further aspect the present invention provides a method of treatment of the human or non-human (preferably mammalian) animal body, said method comprising altering the levels of peptides derived from targeting subunit(s) of PP1c, functional equivalents or portions thereof to an extent that cellular metabolism or function is affected.

Aspects of cellular metabolism that may be affected include (but are not limited to) glycogen metabolism, muscle metabolism, physiology and function.

Generally the levels of peptides or their activity will be enhanced in cells and this control may be achieved by causing higher levels of expression of nucleotides sequences encoding for such peptides (optionally linked to molecules which allow them to cross a cell membrane) or through the administration of such peptides or precursors thereof. Alternatively, in some circumstances, it may be more desirable to depress the levels of certain peptides or at least to depress the level of peptides in active form.

Preferred peptides according to the present invention are derivatives of G_M , especially [G63-T93], [G63-N75], [E2-R575], [E2-P243], [E2-D118], [H100-P350] and peptide 63 to 80, and derivatives of M_{110} , especially

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[M1-E309], [M1-F38], [M1-A150], [L24-Y496]. Preferably, the peptide is not [E2-R575] or [H100-P350].

Particularly preferred peptides are those derived from amino acid nos. 63 to 93 (including 63-80 and 63-75) of G_M ; or from amino acids 1 to 309 (including from 1-150 and 1-38) of M_{110} .

The sequence of G_M is given in Chen et al (1994) Diabetes 43, 1234-1241.

In yet further aspect the present invention provides chimeric proteins containing portions of other proteins or peptides or containing additional amino acids.

Additionally the present invention provides nucleotide sequences (optionally in the form of plasmids) encoding the peptides or chimeric proteins of interest. DNA which encodes the polypeptides or peptides of the invention or chimeric proteins can be made based on a knowledge of the peptide sequences disclosed herein. The DNA is then expressed in a suitable host to produce a polypeptide comprising the compound of the invention. Thus, the DNA encoding the polypeptide constituting the compound of the invention may be used in accordance with known techniques, appropriately modified in view of the teachings contained herein, to construct an expression vector, which is then used to transform an appropriate host cell for the expression and production of the polypeptide of the invention. Such techniques include those disclosed in US Patent Nos. 4,440,859 issued 3 April 1984 to Rutter et al, 4,530,901 issued 23 July 1985 to Weissman, 4,582,800 issued 15 April 1986 to Crowl, 4,677,063 issued 30 June 1987 to Mark et al, 4,678,751 issued 7 July 1987 to Goeddel, 4,704,362 issued 3 November 1987 to Itakura et al, 4,710,463 issued 1 December 1987 to Murray, 4,757,006 issued 12 July 1988 to Toole, Jr. et al, 4,766,075 issued 23 August 1988 to Goeddel et al and 4,810,648 issued 7 March 1989 to Stalker, all of which are incorporated herein by reference.

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The DNA encoding the polypeptide constituting the compound of the invention may be joined to a wide variety of other DNA sequences for introduction into an appropriate host. The companion DNA will depend upon the nature of the host, the manner of the introduction of the DNA into the host, and whether episomal maintenance or integration is desired.

Generally, the DNA is inserted into an expression vector, such as a plasmid, in proper orientation and correct reading frame for expression. If necessary, the DNA may be linked to the appropriate transcriptional and translational regulatory control nucleotide sequences recognised by the desired host, although such controls are generally available in the expression vector. The vector is then introduced into the host through standard techniques. Generally, not all of the hosts will be transformed by the vector. Therefore, it will be necessary to select for transformed host cells. One selection technique involves incorporating into the expression vector a DNA sequence, with any necessary control elements, that codes for a selectable trait in the transformed cell, such as antibiotic resistance. Alternatively, the gene for such selectable trait can be on another vector, which is used to co-transform the desired host cell.

- Host cells that have been transformed by the recombinant DNA of the invention are then cultured for a sufficient time and under appropriate conditions known to those skilled in the art in view of the teachings disclosed herein to permit the expression of the polypeptide, which can then be recovered.
- Many expression systems are known, including bacteria (for example E. coli and Bacillus subtilis), yeasts (for example Saccharomyces cerevisiae), filamentous fungi (for example Aspergillus), plant cells, animal cells and insect cells.
- 30 The vectors include a prokaryotic replicon, such as the ColE1 ori, for

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propagation in a prokaryote, even if the vector is to be used for expression in other, non-prokaryotic, cell types. The vectors can also include an appropriate promoter such as a prokaryotic promoter capable of directing the expression (transcription and translation) of the genes in a bacterial host cell, such as E. coli, transformed therewith.

A promoter is an expression control element formed by a DNA sequence that permits binding of RNA polymerase and transcription to occur. Promoter sequences compatible with exemplary bacterial hosts are typically provided in plasmid vectors containing convenient restriction sites for insertion of a DNA segment of the present invention.

Typical prokaryotic vector plasmids are pUC18, pUC19, pBR322 and pBR329 available from Biorad Laboratories, (Richmond, CA, USA) and pTrc99A and pKK223-3 available from Pharmacia, Piscataway, NJ, USA.

A typical mammalian cell vector plasmid is pSVL available from Pharmacia, Piscataway, NJ, USA. This vector uses the SV40 late promoter to drive expression of cloned genes, the highest level of expression being found in T antigen-producing cells, such as COS-1 cells.

An example of an inducible mammalian expression vector is pMSG, also available from Pharmacia. This vector uses the glucocorticoid-inducible promoter of the mouse mammary tumour virus long terminal repeat to drive expression of the cloned gene.

Useful yeast plasmid vectors are pRS403-406 and pRS413-416 and are generally available from Stratagene Cloning Systems, La Jolla, CA 92037, USA. Plasmids pRS403, pRS404, pRS405 and pRS406 are Yeast Integrating plasmids (YIps) and incorporate the yeast selectable markers HIS3, TRP1,

LEU2 and URA3. Plasmids pRS413-416 are Yeast Centromere plasmids (YCps)

A variety of methods have been developed to operably link DNA to vectors via complementary cohesive termini. For instance, complementary homopolymer tracts can be added to the DNA segment to be inserted to the vector DNA. The vector and DNA segment are then joined by hydrogen bonding between the complementary homopolymeric tails to form recombinant DNA molecules.

Synthetic linkers containing one or more restriction sites provide an alternative method of joining the DNA segment to vectors. The DNA segment, generated by endonuclease restriction digestion as described earlier, is treated with bacteriophage T4 DNA polymerase or *E. coli* DNA polymerase I, enzymes that remove protruding, 3'-single-stranded termini with their 3'-5'-exonucleolytic activities, and fill in recessed 3'-ends with their polymerizing activities.

The combination of these activities therefore generates blunt-ended DNA segments. The blunt-ended segments are then incubated with a large molar excess of linker molecules in the presence of an enzyme that is able to catalyze the ligation of blunt-ended DNA molecules, such as bacteriophage T4 DNA ligase. Thus, the products of the reaction are DNA segments carrying polymeric linker sequences at their ends. These DNA segments are then cleaved with the appropriate restriction enzyme and ligated to an expression vector that has been cleaved with an enzyme that produces termini compatible with those of the DNA segment.

Synthetic linkers containing a variety of restriction endonuclease sites are commercially available from a number of sources including International Biotechnologies Inc, New Haven, CN, USA.

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A desirable way to modify the DNA encoding the polypeptide of the invention is to use the polymerase chain reaction as disclosed by Saiki et al (1988) Science 239, 487-491.

In this method the DNA to be enzymatically amplified is flanked by two specific oligonucleotide primers which themselves become incorporated into the amplified DNA. The said specific primers may contain restriction endonuclease recognition sites which can be used for cloning into expression vectors using methods known in the art. In relation to the above section on DNA expression the term "polypeptide" includes peptides and chimeric proteins.

Further the present invention provides host cells transformed with suitable expression vectors and able to express the peptides. The host cells may be prokaryotic (e.g. E. coli) or eukaryotic (e.g. yeast, mammalian cell cultures).

Bacterial cells are preferred prokaryotic host cells and typically are a strain of *E. coli* such as, for example, the *E. coli* strains DH5 available from Bethesda Research Laboratories Inc., Bethesda, MD, USA, and RR1 available from the American Type Culture Collection (ATCC) of Rockville, MD, USA (No ATCC 31343). Preferred eukaryotic host cells include yeast and mammalian cells, preferably vertebrate cells such as those from a mouse, rat, monkey or human fibroblastic cell line. Yeast host cells include YPH499, YPH500 and YPH501 which are generally available from Stratagene Cloning Systems, La Jolla, CA 92037, USA. Preferred mammalian host cells include Chinese hamster ovary (CHO) cells available from the ATCC as CCL61, NIH Swiss mouse embryo cells NIH/3T3 available from the ATCC as CRL 1658, and monkey kidney-derived COS-1 cells available from the ATCC as CRL 1650.

30 Transformation of appropriate cell hosts with a DNA construct of the present

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invention is accomplished by well known methods that typically depend on the type of vector used. With regard to transformation of prokaryotic host cells, see, for example, Cohen et al (1972) Proc. Natl. Acad. Sci. USA 69, 2110 and Sambrook et al (1989) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY. Transformation of yeast cells is described in Sherman et al (1986) Methods In Yeast Genetics, A Laboratory Manual, Cold Spring Harbor, NY. The method of Beggs (1978) Nature 275, 104-109 is also useful. With regard to vertebrate cells, reagents useful in transfecting such cells, for example calcium phosphate and DEAE-dextran or liposome formulations, are available from Stratagene Cloning Systems, or Life Technologies Inc., Gaithersburg, MD 20877, USA.

Electroporation is also useful for transforming cells and is well known in the art for transforming yeast cell, bacterial cells and vertebrate cells.

For example, many bacterial species may be transformed by the methods described in Luchansky et al (1988) Mol. Microbiol. 2, 637-646 incorporated herein by reference. The greatest number of transformants is consistently recovered following electroporation of the DNA-cell mixture suspended in 2.5X PEB using 6250V per cm at 25μ FD.

Methods for transformation of yeast by electroporation are disclosed in Becker & Guarente (1990) Methods Enzymol. 194, 182.

Successfully transformed cells, ie cells that contain a DNA construct of the present invention, can be identified by well known techniques. For example, cells resulting from the introduction of an expression construct of the present invention can be grown to produce the polypeptide of the invention. Cells can be harvested and lysed and their DNA content examined for the presence of the DNA using a method such as that described by Southern (1975) J. Mol. Biol.

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98, 503 or Berent et al (1985) Biotech. 3, 208. Alternatively, the presence of the protein in the supernatant can be detected using antibodies as described below.

In addition to directly assaying for the presence of recombinant DNA, successful transformation can be confirmed by well known immunological methods when the recombinant DNA is capable of directing the expression of the protein. For example, cells successfully transformed with an expression vector produce proteins displaying appropriate antigenicity. Samples of cells suspected of being transformed are harvested and assayed for the protein using suitable antibodies.

Thus, in addition to the transformed host cells themselves, the present invention also contemplates a culture of those cells, preferably a monoclonal (clonally homogeneous) culture, or a culture derived from a monoclonal culture, in a nutrient medium.

In another aspect the present invention provides antibodies to PP1c which act in an analogous manner to the peptides of interest. Antibodies to the peptides themselves are also provided and these may themselves be used to affect cell metabolism or function.

Peptides in which one or more of the amino acid residues are chemically modified, before or after the peptide is synthesised, may be used providing that the function of the peptide, namely the production of specific antibodies in vivo, remains substantially unchanged. Such modifications include forming salts with acids or bases, especially physiologically acceptable organic or inorganic acids and bases, forming an ester or amide of a terminal carboxyl group, and attaching amino acid protecting groups such as N-t-butoxycarbonyl. Such modifications may protect the peptide from in vivo metabolism. The peptides

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may be present as single copies or as multiples, for example tandem repeats. Such tandem or multiple repeats may be sufficiently antigenic themselves to obviate the use of a carrier. It may be advantageous for the peptide to be formed as a loop, with the N-terminal and C-terminal ends joined together, or to add one or more Cys residues to an end to increase antigenicity and/or to allow disulphide bonds to be formed. If the peptide is covalently linked to a carrier, preferably a polypeptide, then the arrangement is preferably such that the peptide of the invention forms a loop.

According to current immunological theories, a carrier function should be present in any immunogenic formulation in order to stimulate, or enhance stimulation of, the immune system. It is thought that the best carriers embody (or, together with the antigen, create) a T-cell epitope. The peptides may be associated, for example by cross-linking, with a separate carrier, such as serum albumins, myoglobins, bacterial toxoids and keyhole limpet haemocyanin. More recently developed carriers which induce T-cell help in the immune response include the hepatitis-B core antigen (also called the nucleocapsid protein), presumed T-cell epitopes such as Thr-Ala-Ser-Gly-Val-Ala-Glu-Thr-Thr-Asn-Cys (SEQ ID No 1), beta-galactosidase and the 163-171 peptide of interleukin-1. The latter compound may variously be regarded as a carrier or as an adjuvant or as both. Alternatively, several copies of the same or different peptides of the invention may be cross-linked to one another; in this situation there is no separate carrier as such, but a carrier function may be provided by such cross-linking. Suitable cross-linking agents include those listed as such in the Sigma and Pierce catalogues, for example glutaraldehyde, carbodiimide and succinimidyl 4-(N-maleimidomethyl)cyclohexane-1-carboxylate, the latter agent exploiting the -SH group on the C-terminal cysteine residue (if present).

If the peptide is prepared by expression of a suitable nucleotide sequence in a suitable host, then it may be advantageous to express the peptide as a fusion

product with a peptide sequence which acts as a carrier. Kabigen's "Ecosec" system is an example of such an arrangement.

The peptide of the invention may be linked to other antigens to provide a dual effect.

In a yet further aspect the present invention provides a method of diagnosis of abnormalities of cellular metabolism, said method comprising analysing the naturally occurring peptide(s) or the nucleotide sequences encoding therefore and comparing the results to the peptides described herein.

The peptides of the present invention may also be used in diagnosis and this aspect is also covered by the present invention.

The specificity of the catalytic subunit of protein phosphatase-1 (PP1c) is 15 modified by regulatory subunits that target it to particular subcellular locations. For the first time we have identified PP1c-binding domains on G_L and G_M, the subunits that target PP1c to hepatic and muscle glycogen, respectively, and on M₁₁₀, the subunit that targets PP1c to smooth muscle myosin. The peptide G_{M} -(G63-T93) interacted with PP1c and prevented G_{L} from suppressing the 20 dephosphorylation of glycogen phosphorylase, but it did not dissociate G_L from PP1c or affect other characteristic properties of the PP1_{GL} complex. These results indicate that G_L contains two PP1c-binding sites, the region which suppresses the dephosphorylation of glycogen phosphorylase being distinct from that which enhances the dephosphorylation of glycogen synthase. At higher 25 concentrations, G_{M} -(G63-N75) had the same effect as G_{M} -(G63-T93), but not if Ser67 was phosphorylated by cyclic AMP-dependent protein kinase. Thus phosphorylation of Ser67 dissociates G_M from PP1c because phosphate is inserted into the PP1c-binding domain of G_M . The fragments M_{110} -(M1-E309) and M_{110} -(M1-F38), but not M_{110} -(D39-E309), mimicked the M_{110} subunit in 30

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stimulating dephosphorylation of the smooth muscle myosin P-light chain and heavy meromyosin *in vitro*. However, in contrast to the M_{110} subunit and M_{110} -(M1-E309), neither M_{110} -(M1-F38) nor M_{110} -(D39-E309) suppressed the PP1c-catalysed dephosphorylation of glycogen phosphorylase. These observations suggest that the region which stimulates the dephosphorylation of myosin is situated within the N-terminal 38 residues of the M_{110} subunit, while the region which suppresses the dephosphorylation of glycogen phosphorylase requires the presence of at least part of the region 39-296 which contains seven ankyrin repeats. M_{110} -(M1-F38) displaced G_L from PP1c, while G_M -(G63-T93) displaced M_{110} from PP1c *in vitro*. These observations indicate that the region(s) of PP1c that interact with G_M/G_L and M_{110} overlap, explaining why different forms of PP1c contain just a single targeting subunit.

We also disclose the structure of PP1c in complex with a portion of a targeting subunit, and show that changing key amino acid residues in the subunit disrupts its interaction with PP1c. These studies identify a critical structural motif in targeting subunits involved in the interaction with PP1c as well as the recognition site on PP1c itself. These findings will facilitate the rational design of agents such as peptides or other forms of small cell-permeant molecules that act by disrupting PP1-targeting subunit interactions. Given the structural motif and the coordinates of the atoms in the crystal structure, it is within the scope of the abilities of a skilled molecular modeller to produce small cell-permeant molecules, which can enter cells naturally, and possess either the same motif, or an analogous structure to give the same functional properties to the molecule. Thus the small cell-permeant molecule can have a precise copy of the motif, or one which is functionally equivalent. The molecule can be a peptide, but other types of molecules, which are transferred across the plasma membrane of cells, may be preferred.

30 Several mammalian PP1c targeting subunits have been isolated and

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characterised, including the G_M subunit that targets PP1c to both the glycogen particles and sarcoplasmic reticulum of striated muscles (Tang et al., 1991; Chen et al., 1994), the G_L subunit that targets PP1c to liver glycogen (Moorhead et al., 1995; Doherty et al., 1995), the M₁₁₀ subunits responsible for the association of PP1c with the myofibrils of skeletal muscle (Moorhead et al., 1994; Alessi et al., 1992) and smooth muscle (Alessi et al., 1992; Chen et al., 1994), the p53 binding protein p53BP2 (Helps et al., 1995) and the nuclear protein NIPP-1 (Jagiello et al., 1995; Van Eynde et al., 1995). PP1c is also reported to interact with other mammalian proteins such as the retinoblastoma gene product (Durphee et al., 1993), an RNA splicing factor (Hirano et al., 1996), ribosomal proteins L5 (Hirano et al., 1995) and RIPP-1 (Beullens et al., 1996), a 110 kDa nuclear protein yet to be identified (Jagiello et al., 1995), kinesin-like proteins and small cytosolic proteins, inhibitor-1, DARPP-32 and inhibitor-2 (Cohen, 1989; Cohen, 1992, Hubbard and Cohen, 1993). Moreover, a number of distinct PP1-regulatory subunits have been identified in yeast (reviewed by Stark, 1996). We attempted to identify which regions of the G_M and M_{110} subunits were involved in binding to PP1c. These studies led to the identification of relatively small peptides from each targeting subunit that were capable of interacting with PP1c. Peptides comprising residues 63-93, 63-80 and 63-75 of G_M bound to PP1c, dissociating it from G_L, while the N-terminal 38 residues of the M_{110} subunit ($M_{110}[1-38]$) mimicked the intact M₁₁₀ subunit in enhancing the rate at which PP1c dephosphorylated the 20 kDa myosin light chain (MLC₂₀) subunit of smooth muscle myosin (Johnson et al., 1996).

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The present invention thus provides peptides comprising the N-terminal 38 residues of the M_{110} subunit, and those comprising residues 63-93, 63-80 and 63-75 of G_M .

Phosphorylation of Ser 67 of G_M by protein kinase A (PKA) disrupts the

interaction of G_M with PP1c (Dent et al., 1990) and a similar disruption is also observed following the phosphorylation of Ser 67 of the $G_M[63-75]$ peptide (Johnson et al., 1996). The finding that $G_M[63-93]$ disrupted the interaction between PP1c and the M_{110} subunit, and prevented M_{110} from enhancing the MLC_{20} phosphatase activity of PP1c implies that the binding of M_{110} and G_M to PP1c are mutually exclusive.

Thus the invention contemplates the substitution or modification of an amino acid in any such peptide.

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To understand the basis for the recognition by PP1c of regulatory subunits, and peptides derived from these subunits, we co-crystallised a complex of PP1c with the $G_M[63-75]$ peptide and determined the structure at 3.0 Å resolution. These experiments have demonstrated that residues 64 to 69 of the peptide are bound in an extended conformation to a hydrophobic channel within the C-terminal region of PP1c. The residues in $G_M[63-75]$ that interact with PP1c lie in an Arg/Lys-Val/Ile--Xaa-Phe motif common to $M_{110}[1-38]$ and almost all known mammalian PP1-binding proteins. Substituting Val or Phe by Ala in the $G_M[63-75]$ peptide, and deleting the VxF motif from the $M_{110}[1-38]$ peptide, abolished the ability of both peptides to interact with PP1c. These findings identify a recognition site on PP1c for a critical structural motif involved in the interaction of targeting subunits with PP1.

Particularly preferred peptides are derived from residues 63 to 69 of G_M and comprise the motif Arg/Lys-Val/Ile-Xaa-Phe. Peptides derived from M_{110} (or any other source) and also including the motif are also included in the scope of the invention.

Preferred peptides may also be substantially or wholly made up of hydrophobic residues.

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The identification of this area of PP1c necessary for binding to the various subunits allows the design of agents to specifically disrupt the interaction at this area. Such disruption may, for example, increase the phosphorylation of the protein phospholamban in cardiac muscle and thus increase the force and rate of contraction of the muscle. This provides a possible treatment for congestive heart failure. Also, the specific disruption of the complex of PP1 and p53BP2 may prevent PP1 from dephosphorylating the tumour suppressor protein p53, thus enhancing phosphorylation of p53, its ability to bind to DNA, and thus its ability to act as a tumour suppressor.

The identification of the key motif in targetting subunits that bind to PP1 also provides the means to produce targetting subunits that can no longer interact with PP1. Over-expression of these mutant targetting subunits provides a powerful new way to determine the functions of different targetting subunits in vivo.

Abbreviations

20 PP1, protein phosphatase-1

PP1c, catalytic subunit of PP1

PP1, -isoform of PP1c

PP1_G, glycogen-associated form of PP1

PP1_M, myosin-associated form of PP1

25 G_M, glycogen-binding subunit of PP1 from striated muscle

G_L, glycogen-binding subunit of PP1 from liver

NIPP1, nuclear inhibitor of PP1

DARPP, dopamine and cyclic AMP-regulated phosphoprotein

 M_{21} and M_{110} , myofibrillar-binding subunits of PP1 with molecular masses of

30 21kDa and 110 kDa respectively.

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PKA, cyclic AMP-dependent protein kinase
PhMeSO₂F, phenylmethylsuphonyl fluoride
GST, glutathione-S-transferase
MLC₂₀, myosin light chain of molecular mass 20 kDa.

The invention is now described in more detail by reference to the following Examples and Figures wherein:

Figure 1 shows that the N-terminal 118 residues of human G_M interact with PP1c.

GST-G_M fusion proteins were electrophoresed on 10% SDS/polyacrylamide gels and stained with Coomassie blue (lanes 1-3) or, after transfer to nitrocellulose, probed with digoxygenin-labelled PP1γ (lanes 4-6) as in [9]. Lanes 1 and 4, GST-G_M-(E2-D118); Lanes 2 and 5, GST-G_M-(H100-P350); Lanes 3 and 6, GST. The positions of the marker proteins glycogen phosphorylase (97 kDa), bovine serum albumin (66 kDa), ovalbumin (43 kDa), carbonic anhydrase (29 kDa) and soybean trypsin inhibitor (20 kDa) are indicated.

Figure 2 shows that synthetic peptides between residues 63 and 93 of rabbit G_M stimulate the phosphorylase phosphatase activity of PP1_{GL}.

Hepatic glycogen particles were diluted in assay buffer to 0.6 phosphorylase phosphatase (PhP) mU per ml, incubated for 15 minutes at 30°C with G_{M} -(G63-T93) (closed circles), G_{M} -(G63-K80) (open circles) or G_{M} -(G63-N75) (closed triangles) and assayed as described in Example 1. The open triangles show the effect of G_{M} -(G63-N75) which had been phosphorylated at Ser67 by PKA (p G_{M} -(G63-N75)). Similar results were obtained in four experiments.

Figure 3 shows that removal of the M_{21} subunit from smooth muscle $PP1_M$ does

not affect its MLC₂₀ phosphatase:phosphorylase phosphatase activity ratio.

- (A) Purified smooth muscle $PP1_M$ was electrophoresed on a 12% SDS/polyacrylamide gel, and either stained with Coomassie blue (lane 1) or immunoblotted [32] with antibodies specific for the M_{21} subunit (lane 2) or the M_{110} subunit (lane 3). The positions of the M_{110} subunit, the M_{21} subunit and PP1c are marked.
- (B) Purified PP1_M (lane 1) or PP1_M lacking the M₂₁ subunit (lane 2) were electrophoresed on a 12% SDS polyacrylamide gel, transferred to nitrocellulose and immunoblotted with mixed, affinity-purified antibodies to the M₁₁₀ and M₂₁ subunits. The M₁₁₀ and M₂₁ subunits are marked. The activity ratio, MLC₂₀ phosphatase (MP):phosphorylase phosphatase (PhP) of the two preparations is also shown. Similar results were obtained in three different experiments. The activity ratio MP:PhP of PP1c is 0.07.
 - Figure 4 shows expressed fragments of the M_{110} subunit before and after cleavage of the GST-fusion proteins with thrombin .
- electrophoresed Purified **GST-fusion** proteins were 15% 20 SDS/polyacrylamide gel and stained with Coomassie blue. Lane 1, E309); $GST-M_{110}-(M1-A150);$ $GST-M_{110}-(D39-$ 3, Lane 2, GST- M_{110} -(M1-E309); Lane 4, GST- M_{110} -(L24- Y496). Lanes 5-8 are the same as Lanes 1-4 except that the fusion proteins were cleaved with thrombin. The positions of the marker proteins glycogen phosphorylase (97 kDa), bovine 25 serum albumin (66 kDa), ovalbumin (43 kDa), carbonic anhydrase (29 kDa), GST (26 kDa) and soybean trypsin inhibitor (20 kDa) are marked.
- Figure 5 shows the effect of M₁₁₀ subunit fragments on PP1c-catalysed dephosphorylation of MLC₂₀ and glycogen phosphorylase

A,B; Effects of M_{110} -(M1-E309) (closed circles), M_{110} -(M1-F38) (open circles) and M_{110} -(D39-E309) (open triangles) on the MLC₂₀ phosphatase (B) and phosphorylase phosphatase (B) activities of PP1c were measured after incubating PP1c for 15 minutes at 30°C with each fragment. The results are presented as a percentage of those obtained in experiments where the M_{110} fragments were omitted.

C,D; The effect of M_{110} -(M1-A150) (open circles) and M_{110} -(L24-Y496) (closed circles) on the MLC₂₀ phosphatase (C) and phosphorylase phosphatase (D) activities of PP1c were measured as in A,B.

Figure 6 shows the effect of M_{110} -(M1-F38) and M_{110} -(M1-E309) on the dephosphorylation of glycogen synthase by PP1c.

- The glycogen synthase phosphatase activity of PP1c was measured after a 15 minute incubation at 30°C with the indicated concentrations of M₁₁₀-(M1-F38) and M₁₁₀-(M1-E309). Similar results were obtained in three different experiments.
- 20 Figure 7 shows that G_M-(G63-T93) dissociates PP1_M.
- (A) The phosphorylase phosphatase (PhP) activity of PP1_M (closed circles) and its MLC₂₀ phosphatase (MLCP) activity (open circles) were assayed after preincubation for 15 minutes at 30°C with the indicated concentrations of G_M-(G63-T93). Activities are shown relative to control incubations in which G_M-(G63-T93) was omitted. Similar results were obtained in three experiments.
- (B,C) PPIM was incubated for 15 minutes at 30°C in the absence (B) and presence (C) of 10 M G_M-(G63-T93), then passed through a 30 x 1 cm column

of Superose 12 equilibrated at ambient temperature in 50 mM Tris/HC1 pH 7.5, 0.2M NaC1, 0.1 mM EGTA, 0.1% (by vol) 2-mercaptoethanol, 0.03% (by mass) Brij 35 in the absence (B) or presence (C) of 1 μ M G_M-(G63-T93). Fractions (0.25 ml) were assayed for MLC₂₀ phosphatase (MLCP) in B and for phosphorylase phosphatase (PhP) activity in C. The arrows denote the position of ferritin (450 kDa) and ovalbumin (43 kDa).

Figure 8 shows that G_{M} -(G63-T93) prevents M_{110} -(M1-F38) or M_{110} -(M1-E309) from modulating the substrate specificity of PP1c.

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- (A) The MLC₂₀ phosphatase activity of PP1c was assayed after incubation for 15 minutes at 30°C in the presence or absence of 1 μ M G_M-(G63-T93) and either 0.1 μ M M₁₁₀-(M1-F38) or 0.1 nM M₁₁₀-(M1-E309).
- (B) The phosphorylase phosphatase activity of PP1c was assayed as in A in the presence or absence of 1 μM G_M-(G63-T93) and 1.0 nM M₁₁₀-(M1-E309). The results are presented (SEM for three experiments) as a percentage of the PP1c activity measured in the absence of G_M-(63-T93), M₁₁₀-(M1-F38) or M₁₁₀-(M1-E309).

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Figure 9 shows the location of PP1c-binding domains on the G_M and M_{110} targeting subunits and their effects on PP1 activity.

The hatched boxes in the M_{110} subunit denote the positions of the ankyrin repeats.

Figure 10 shows a stereo view of the electron density corresponding to the peptide. A: Initial 2-fold averaged electron map. B: map calculated using 3Fo-2Fc coefficients and phases calculated from the final refined model. Displayed using TURBO-FRODO.

Figure 11 shows the structure of PP1- $G_M[63-75]$ peptide complex. A. Stereo view of a ribbons diagram of PP1c to indicate the position of the peptide binding channel at the interface of the two β -sheets of the β -sandwich. The peptide atoms are represented as ball-and-stick (MOLSCRIPT, Kraulis, 1991).

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- B. View of the surface of PP1c to show the hydrophobic peptide binding channel. Residues 63' to 69' (GRRVSFA) (SEQ ID No 2) of the G_M [63 75] peptide are shown as sticks. Drawn with TURBO-FRODO.
- 10 C. Stereo view of the $G_M[63-75]$ peptide at the recognition site of PP1 to indicate polar interactions between peptide and protein and the formation of the β -sheet between Ser 67' Ala 69' and 14 of PP1. Drawn with TURBO-FRODO.
- D. Solvent accessible surface and surface electrostatic potential of PP1-G_M[63 75] peptide complex calculated with PP1 coordinates alone and showing the peptide as a stick representation in the vicinity of the peptide binding site. The figure was created with GRASP (Nicholls and Honig, 1991). The protein surface is coloured according to electrostatic potential from red (most negative) to blue (most positive). The figure shows pronounced negative electrostatic potential in the region surrounding the N-terminus of the peptide binding site that results from seven conserved acidic residues.
- E. Details of the structure of the peptide binding site to show hydrophobic interactions between PP1c and Val 66', Phe 68' and Ala 69' of the G_M[68-75] peptide (MOLSCRIPT, Kraulis, 1991).

Figure 12 shows a sequence alignment of PP1-regulatory subunits in the vicinity of the (R/K)(V/I) x F motif. (A) mammalian PP1-binding subunits.

30 G_M, Tang et al., 1991; GL, Docherty et al., 1995; G_L-related protein, Doherty

et al., 1996; p53BP2, Helps et al., 1995; NIPP-1, Bollen et al., 1995; splicing factor PSF, Hirano et al., 1996; M₁₁₀ subunit, Chen et al., 1994; inhibitor-1, Aitken et al., 1982; DARPP-32, Williams et al., 1986. (B) PP1-binding proteins in S. cerevisiae. GACl (Francois et al., 1992); PIG2 GIP1, GIP2, YILO45W (Tu et al., 1996); REG1, REG2 (Tu and Carlson, 1995; Frederick and Tatchell, 1996); SCD5 (Nelson et al 1996; Tu et al 1996). The region homologous to the RRVSFA (SEQ ID No 3) motif in G_M which intersects with PP1c is boxed.

Figure 13 shows the disruption of the interactions between PP1c and the G_L and M₁₁₀ subunits by a synthetic peptide from p53BP2. (A) PP1_M from chicken gizzard smooth muscle (Alessi et al., 1992) was diluted and incubated for 15 m i n 3 0 ° C with t h e GKRTNLRKTGSERIAHGMRVKFNPLALLLDSC (SEQ ID No corresponding to the sequence in p53BP2 that contains the RVxF motif. Reactions were started with either ³²P-labelled MLC₂₀ or glycogen phosphorylase and the MLC₂₀ phosphatase (open circles) and phosphorylase phosphatase (PhP, closed circles) activities were determined. The results are expressed as a percentage of the activity determined in control incubations where the p53BP2 peptide was omitted (100%). Similar results were obtained in three separate experiments. (B) same as (A) except that the peptide was incubated with diluted hepatic glycogen particles containing PP1-G_L before measuring the PhP activity. Similar results were obtained in three separate experiments.

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Figure 14 shows the effect of $M_{110}[M1-F38]$ and $M_{110}[M1-K35]$ on the PP1c-catalysed dephosphorylation of MLC₂₀ $M_{110}[M1-F38]$ (1-38, open circles) or $M_{110}[M1-K35]$ (1-35, closed circles) were incubated with PP1c for 15 min at 30°C and reactions started with the ³²P-labelled MLC₂₀ substrate. The results are expressed as a % of the activity determined in control incubations where

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the M_{110} peptides were omitted (100%). Similar results were obtained in three separate experiments.

Figure 15 shows the effect of synthetic peptides derived from the M_{110} and G_M subunits on the phosphorylase phosphatase activity of PP1- G_L . (A) Hepatic glycogen protein particles containing PP1- G_L were diluted and incubated for 15 min at 30°C with the indicated concentrations of either $M_{110}[M1-F38]$ (open circles) or $M_{110}[M1-K35]$ (closed circles) and the phosphatase reactions were initiated by addition of ^{32}P -labelled glycogen phosphorylase. The results are expressed as a percentage of the activity determined in control incubations where the M_{110} peptides were omitted. Similar results were obtained in three separate experiments. (B) The experiment was carried out as in (A), except that the peptide $G_M[G63-N75]$ ("wild type", WT) and variants in which either Val 66 (V66A) (closed triangles) or Phe 68 (F68A) (closed circles) were changed to Ala, were used instead of the M_{110} peptides. Similar results were obtained in three separate experiments.

Figure 16 shows a stereo view of a ribbons diagram of a model of PP1-phospho-inhibitor-l complex. The side chains of Ile 10, Phe 12 and pThr 35 of phospho-inhibitor-l are shown with the main-chain atoms of residues 8 to 36 of the inhibitor indicated as a shaded ribbon. Drawn with MOLSCRIPT (Kraulis 1991).

Figure 17 shows a comparison of rat and chicken gizzard M_{110} and M_{21} subunits.

Vertical lines indicate identical residues, colons denote similar residues in the rat and chicken M_{110} sequences and deletions are shown by dots. (A) Comparison of M_{110} subunits. Underlined residues in the rat M_{110} subunit (Rat1) are deleted in some rat aorta forms and underlined residues in the

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chicken M_{110} subunit (Ch1) are deleted in some chicken gizzard forms [5, 8]. Dashed lines above residues indicate amino acids deleted in the rat kidney M_{110} subunit [9]. The alternative C-terminal sequences of rat uterus M_{110} subunit are shown as Rat1 and Rat2. Leucine residues in the C-terminal leucine zipper motif are double underlined. (B). The C-terminal sequence of the M_{110} subunit is structurally related to the M_{21} subunit. The sequence of the chicken M_{21} subunit [5] is compared with the C-terminal sequences of Rat2 and Ch1 from A. Identities between Ch1 and Rat2 are shown in boldface type.

Figure 18 shows immunoprecipitation and immunoblotting of PP1_M in extracts from chicken gizzard myofibrils.

A. Antibodies specific for the M_{110} and/or M_{21} subunits immunoprecipitate most of the myosin P-light chain phosphatase activity in myofibrillar extracts. PP1_M was immunoprecipitated with either control IgG, antibody raised against the PP1_M holoenzyme, antibody specific for the M₁₁₀ subunit or antibody specific for the M₂₁ subunit, as described under Methods in Example 3. The figure shows activity present in the supernatant (S, open bars) or pellet (P, filled bars) as a percentage of that measured before centrifugation. The results shown are the average (± S.E.M.) for three separate experiments each assayed in duplicate. B, The M_{110} and M_{21} subunits are present in similar molar proportions in myofibrillar extracts and in purified PP1_M. 10 ng (track 1) or 3 ng (track 3) of purified PP1_M or 12 μ g (track 2) or 3.6 μ g (track 4) of myofibrillar extract was electrophoresed on a 12% SDS/polyacrylamide gel, transferred to nitrocellulose and immunoblotted with mixed affinity-purified antibodies to the M_{110} and M_{21} subunits as in [22]. The positions of the two subunits are marked. The results indicate that PP1_M comprises about 0.1% of the myofibrillar protein.

30 Figure 19 shows the identification of the region on the M₁₁₀ subunit that

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interacts with the M21 subunit.

A) PP1_M 5 μ g (track 1), 10 μ g bacterial extract containing M₁₁₀-(R714-I1004) (track 2), MBP-M₁₁₀-(R714-I1004) 1 μ g (track 3), MBP-M₁₁₀-(R714-L934) 1 μ g (track 4), MBP-M₁₁₀-(K933-I1004) 1 μ g (track 5), MBP 1 μ g (track 6), M₁₁₀-(M1-E309) 2 μ g (lane 7) and M₁₁₀-(M1-S477) 2 μ g (track 8) were run on a 12% SDS/polyacrylamide gel and stained with Coomassie Blue. B) same as A) except that 10-fold less protein was electrophoresed and after transfer to nitrocellulose the proteins were probed with digoxigenin-labelled M₂₁ subunit (0.2 μ g/ml). C) same as B) except that, after electrophoresis, the proteins were transferred to nitrocellulose and probed with digoxigenin-labelled M₂₁-(M1-L146) (0.2 μ g/ml).

Figure 20 shows the identification of the region of the M_{21} subunit involved in interaction with the M_{110} subunit and in dimerization.

A) GST-M₂₁ 5 μ g (track 1), M₂₁ 5 μ g (track 2), M₂₁-(M1-L146) 5 μ g (track 3), M₂₁-(M1-E110) 20 μ g (track 4) and M₂₁-(E110-K186) 5 μ g (track 5) were run on 16.5% polyacrylamide gels and stained with Coomassie Blue. The marker proteins ovalbumin (43 kDa) and carbonic anhydrase (29 kDa) are indicated.

B) GST-M₂₁ 0.5 μ g (track 1), M₂₁ 0.5 μ g (track 2), M₂₁-(M1-L146) 0.5 μ g (track 3), M₂₁-(M1-E110) 5 μ g (track 4) and M₂₁-(E110-K186) 5 μ g (track 5) were electrophoresed as in A) and after transfer to nitrocellulose the blots were probed with digoxigenin-labelled MBP-M₁₁₀-(K933-I1004) (0.2 μ g/ml). C) same as B) except that, after electrophoresis, the proteins were transferred to nitrocellulose and probed with digoxigenin-labelled M₂₁ subunit (0.2 μ g/ml).

Figure 21 shows that the M_{21} subunit and M_{21} -(M1-L146) interact with the M_{110} subunit and themselves, but not with PP1.

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PP1_M (0.5 μ g) was electrophoresed on a 12% SDS/polyacrylamide gel, transferred to nitrocellulose and probed with digoxigenin-labelled M₂₁ subunit (0.2 μ g/ml) (track 1) or digoxigenin-labelled M₂₁-(M1-L146) (0.2 μ g/ml) (track 2). The positions of the M₁₁₀ subunit, the M₂₁ subunit and PP1c are marked.

Figure 22 shows that removal of the M_{21} subunit from smooth muscle $PP1_M$ does not prevent it from being pelleted with myosin.

The PP1 catalytic subunit (PP1c), PP1_M, or PP1_M lacking the M_{21} subunit, PP1_M(ΔM_{21}), each at 30 nM, were incubated for 15 min at 0°C with 1 μ M myosin and centrifuged (see Methods of Example 3). The figure shows the myosin P-light chain phosphatase activity present in the supernatant (S, open bars) or pellet (P, filled bars) as a percentage of that measured before centrifugation. The results shown are the average (\pm S.E.M.) for three separate experiments each assayed in duplicate.

Figure 23 shows the identification of a region of the M_{110} subunit which binds to myosin.

(A); PP1_M, M₁₁₀-(M1-S477) and GST-M₁₁₀-(M377-K976), each at 30 nM were incubated for 15 min at 0°C with 1 μM myosin and centrifuged. The supernatants (S), resuspended pellets (P) and the suspension before centrifugation (T, total) were electrophoresed on 12% SDS/polyacrylamide gels, transferred to nitrocellulose and immunoblotted with antibodies raised against the PP1_M holoenzyme. No protein was pelleted in the absence of myosin (not shown). The positions of the marker proteins myosin heavy chain (200 kDa), glycogen phosphorylase (97 kDa), bovine serum albumin (66 kDa), ovalbumin (43 kDa), carbonic anhydrase (29 kDa) and soybean trypsin inhibitor (20 kDa) are indicated. (B) The experiments were carried out as in (A), except that the M₁₁₀ fragments and M₂₁ subunit were used at 100 nM, the 8.5 kDa

 M_{110} -(K933-I1004) fragment was electrophoresed on a 16.5% polyacrylamide gel and immunoblotting was carried out with affinity purified antibodies (see Methods). A small amount of M_{110} -(R714-I1004) pelleted in the absence of myosin. This was probably due to aggregation in the bacterial extract since this did not happen when it was complexed to the M_{21} subunit (data not shown). No other protein was pelleted in the absence of myosin.

Figure 24 shows that the isolated M₂₁ subunit binds to myosin.

- (A); Myosin (1 μM) was mixed with 50 μM, 20 μM or 10 μM M₂₁ subunit to give the molar ratios M₂₁:myosin dimer indicated. After 15 min at 0°C, the solutions were centrifuged and the supernatants (S), resuspended pellets (P) and the suspension before centrifugation (T, total) were electrophoresed on 12% SDS/polyacrylamide gels and stained with Coomassie blue. The positions of the myosin heavy chain (MHC) and the M₂₁ subunit are indicated. The myosin light chains migrate faster than the M₂₁ subunit and are not visible at these loadings.
- (B); Myosin (track A) was purified from chicken gizzard, and the myosin "rod" domain (track B) and light meromyosin (track C) produced by digestion 20 of myosin with papain and chymotrypsin, respectively. These three proteins, all at 1 μ M, were then mixed with M_{21} subunit (track D) to give a molar ratio M₂₁:myosin dimer of 10:1 and, after 15 min at 0°C, the solutions were centrifuged and the supernatants (S), resuspended pellets (P) and the suspension 25 before centrifugation (T, total) were electrophoresed 12% SDS/polyacrylamide gels and stained with Coomassie blue. The slightly faster migrating band in the M₂₁ subunit preparation was shown by amino acid sequencing to be N-terminally truncated commencing at residue 16. (C); same as (B), except that M_{21} -(M1-L146) (track D) replaced the M_{21} subunit.

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Figure 25 gives a schematic representation of the regions on the M_{110} subunit from chicken gizzard that interact with PP1c, myosin and the M_{21} subunit.

PP1c binds to the KVKF (SEQ ID No 5) motif between residue 35 and 38, just N-terminal to the seven ankyrin repeats (hatched vertical lines) that suppress the dephosphorylation of substrates other than myosin. Residues 1-38 of the M_{110} subunit enhance the dephosphorylation of myosin. The M_{21} subunit binds to the C-terminal 72 residues of the M_{110} subunit which are 43% identical in amino acid sequence to residues 87-161 of the M_{21} subunit. The dephosphorylated form of myosin binds to M_{110} -(R714-I1004) but not to M_{110} -(K933-I1004), suggesting that myosin binds N-terminal to the M_{21} subunit.

Example 1: Identification of protein phosphatase 1-binding domains on the glycogen and myofibrillar targeting subunits

MATERIALS AND METHODS

Materials.

The myosin-associated form of PP1 (PP1_M) was from chicken gizzard [9] and the glycogen-associated form of PP1 (PP1_G) from rabbit skeletal muscle [21]. The β isoform of PP1c was released from PP1_G by incubation for 2 hours in 2M LiBr, then purified by gel-filtration on a 30 x 1 cm column of Superose 12 (Pharmacia, Milton Keynes, U.K.) in the presence of 0.5M LiBr. Glycogen protein particles from rat liver [22] were used as the source of hepatic PP1_G. Digoxygenin-labelled PP1c (γ_1 -isoform, hereafter termed PP1) was prepared as in [9]. G_L was expressed in E. coli as a glutathione-S-transferase (GST) fusion protein [7], termed GST-G_L. The catalytic subunit of PP2A from bovine heart (PP2AC) was provided by Dr R. MacKintosh in this Unit. The phosphorylatable myosin light chain (MLC₂₀) and heavy meromyosin from chicken gizzard were a gift from Dr M. Ikebe (Case Western Reserve

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University, Cleveland, USA). Thrombin and benzamidine-Agarose were purchased from Sigma (Poole, UK).

Peptide synthesis.

Peptides were synthesised on an Applied Biosystems 430A peptide synthesiser and their purity and concentration established by high performance liquid chromatography, mass spectrometry and amino acid analysis. The sequence of rabbit G_M-(G63-T93) is GRRVSFADNFGFNLVSVKEFDTWELPSVSTT (SEQ IDNo 6) and the sequence of M_{110} -(M1-F38) MKMADAKQKRNEQLKRWIGSETDLEPPVVKRQKTKVKF (SEQ ID No 10 The peptide G_M-(G63-T93) was cleaved with Lys-C endoproteinase (Boehringer) and the peptide G_M-(E81-T93) thus generated was purified on a The peptides G_{M} -(G63-K80) and G_{M} -(G63-N75), were synthesised, and the latter phosphorylated at Ser67 with the catalytic subunit of cyclic AMP-dependent protein kinase (PKA), then bound to a 1 ml C₁₈ column 15 equilibrated in 0.1% (v/v) trifluoroacetic acid, washed with 0.1% trifluoroacetic acid to remove excess ATP, eluted with 0.1% trifluoroacetic acid containing 70% acetonitrile, dried and dissolved in water. G_M-(S40-Y55) was a gift from Dr Bruce Kemp (St Vincent's Institute, 20 Australia).

Preparation of phosphorylated proteins and phosphatase assays.

³²P-labelled rabbit skeletal muscle phosphorylase a (containing 1.0 mol phosphate per mol subunit) was prepared by phosphorylation with phosphorylase kinase [23], ³²P-labelled rabbit skeletal muscle glycogen synthase (containing 1.5 mol/mol subunit in the sites 3 region) was prepared by phosphorylation with glycogen synthase kinase-3 [24]), ³²P-labelled chicken gizzard MLC₂₀ and ³²P-labelled chicken gizzard heavy meromyosin (containing 1.0 mol phosphate per mol subunit) were prepared by phosphorylation with smooth muscle myosin light chain kinase [9]. The dephosphorylation of

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phosphorylase a (10 μ M), glycogen synthase (1 μ M) and MLC₂₀ (1 μ M) and heavy meromyosin (1 μ M) was carried out as in [24]. One unit of activity (U) was that amount which released 1 mole of phosphate in one minute.

Construction of vectors for the expression of N-terminal fragments of the G_M subunit as glutathione-S-transferase (GST) fusion proteins in E. coli.

 G_{M} -(E2-R575) was produced by inserting a SmaI-SmaI restriction fragment, encoding amino acids 2-575 of human G_{M} , from clone H1G11 [5] into the SmaI site of pGEX-KG (Pharmacia, Milton Keynes, U.K.). This resulted in the addition after residues 2-575 of amino acids EFPVVVVEF (SEQ ID No 8) before the stop codon. G_{M} -(E2-P243) was made by deleting an NcoI-HindIII fragment of the G_{M} -(E2-R575) construct, resulting in termination after residue 243. G_{M} -(E2-D118), encoding amino acids 2-118, with a C-terminal addition of QLNSS was produced by deleting a BgIII-HindIII fragment of the G_{M} -(E2-R575) construct. G_{M} -(H100-P350) encoding amino acids 100-350 was made by inserting an EcoRI-HindIII digested PCR fragment prepared using primers

5' GCCGAATTCACACAGAAGAATATGTTTTAGCC 3' (SEQ ID No 9) and 5' GCCGAAGCTTATGGAAAATTGACTGGATCTGTTG 3' (SEQ ID No 10) into the same sites of pGEX-KG. Restriction sites in the primers are underlined.

Construction of vectors for the expression of the chicken gizzard M_{21} subunit in E. coli.

The entire coding region (M1-K186) of the M_{21} subunit [10] was amplified by PCR using primers

- 5' CGCGCATATGTCGTCGCTGTTCACCAGG 3' (SEQ ID No 11) and
- 30 5' GGCGGATCCCTACTTGGAGAGTTTGC 3' (SEQ ID No 12), containing

restriction sites NdeI and BamH1 (underlined). After cleavage with the restriction enzymes, the PCR fragment was cloned into the same sites of the bacterial expression vector pT7-7.

5 Production of fragments of the chicken gizzard and rat aorta M_{110} subunits.

The C-terminal 291 residues M_{110} -(R714-I1004) of the chicken gizzard M_{110} subunit were amplified by PCR using a primer

5' AGGAAGAATTCGTTCCACACGAAC 3' (SEQ ID No 13) containing an EcoRI restriction site (underlined) and a KS primer in the Bluescript vector of the cDNA clone [10]. The EcoRI digested PCR fragment was subcloned into the same site of pT7-7.

Rat aorta M_{110} fragments were produced as GST-fusion proteins. M₁₁₀-(M1-A150) was amplified by PCR using primers A 15 (5' CCTAGCCCGGGGATGAAGATGGCGGAC 3') (SEQ ID No 14) and B (5' GCGGAAGCTTATGCTTCCTCCTCTGCAATATC 3') (SEQ ID No 15). containing Smal and HindIII restriction sites (underlined) and the Smal-HindIII digested PCR fragment subcloned into the same sites of pGEX-KG. M₁₁₀-(M1-E309) was produced by subcloning a Smal-HindIII digested PCR 20 fragment amplified using primers Α and CTAGAAGCTTCCATATTTGCTGTTGATTCAATC 3') (SEQ ID No 16) into the same sites of pGEX-KG. This resulted in one amino acid (A) being added after E309. M₁₁₀-(D39-E309) was produced by subcloning a Smal-HindIII 25 digested **PCR** fragment amplified D using primers (5' CCTAGCCCGGGGACGATGGCGCCGTCTTCC 3') (SEQ ID No 17) and C into the same sites of pGEX-KG. An M_{110} -(L24-K976) was prepared by inserting a Xhol-Xhol restriction fragment of the entire M₁₁₀ cDNA in Bluescript into XhoI site of pGEX-KG, and M₁₁₀-(L24-Y496) expressed by deleting a Ndel-Ndel fragment of the L24-K976 construct and filling the 30

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overhanging ends before ligating them. This resulted in the addition after Y496 of amino acids MVAD (SEQ ID No 18) before the stop codon. The sequence of all subclones produced after PCR amplification were verified using an Applied Biosystems 373A automated DNA sequencer and Taq dye terminator cycle sequencing according to the manufacturer's instructions.

Expression of proteins in E.coli.

All constructs were expressed in *E. coli* strain BL21(DE3)plysS. Cultures were grown at 37°C in Luria-Bertani medium in the presence of 100 μg/ml ampicillin and 30 μg/ml chloramphenicol to an A600 of 0.4-0.6, and induced with 50 μg/ml isopropylthiogalactoside for 8 hours at 25°C or overnight at ambient temperature. After centrifugation for 10 minutes at 7000 x g (4°C), cells from one litre of culture were resuspended in 20 ml of 50 mM Tris-HCl pH 8.0, 0.1 M NaCl, 1 mM EDTA, 0.1% (by vol) 2-mercaptoethanol, 0.2 mM phenylmethylsulphonylfluoride (PhMeSO₂F), 1 mM benzamidine (buffer A) and frozen at -80°C. After thawing, sodium deoxycholate (1 mg/ml), 8 mM MgSO4 and 10 g/ml DNAase I were added, the extract incubated until it was no longer viscous, then made 6 mM in EDTA, 1 mM in benzamidine and 0.2 mM in PhMeSO₂F and centrifuged for 10 minutes at 10,000 x g. The soluble GST-fusion proteins were then purified from the supernatant by affinity chromatography on glutathione-Sepharose (Pharmacia).

The M_{21} subunit and M_{110} -(R714-I1004) C-terminal fragment from chicken gizzard M_{110} subunit, which were used for affinity purification of the anti- M_{21} and anti- M_{110} antibodies (see below) were obtained in inclusion bodies and therefore recovered in the pellets after centrifuging E. coli extracts at 10,000 x g. M_{110} -(R714-I1004) was solubilised by resuspension in Buffer A containing 0.5% (by mass) Triton X-100 and was >95% pure. The M_{21} subunit was not solubilised by this procedure but, after washing the pellets in 0.5% Triton

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X-100, was dissolved by sonication in 0.5% trifluoroacetic acid; its purity was about 20%.

M₁₁₀ GST-fusion proteins (1-9 mg/ml in 50 mM Tris/HCl, 2.5 mM CaCl2, 150 mM NaCl and 0.1% (by vol) 2-mercaptoethanol) were cleaved by incubation for 20 minutes at 30°C with 20 μg/ml thrombin. Benzamidine-Agarose (0.2 ml) was added and, after incubation (with rotation) for 30 minutes at ambient temperature, the benzamidine-Agarose containing the attached thrombin was removed, and the supernatant dialysed against 50 mM Tris-HCl pH 7.5, 0.1 mM EGTA, 0.1% (by vol) 2-mercaptoethanol, 10% glycerol and stored in aliquots at -80°C. After cleavage with thrombin, all fragments of the M₁₁₀ subunit, except M₁₁₀-(L24-Y496), commenced with the sequence GSPG (SEQ ID No 19) before the initiating residue of the GST-fusion proteins. The M₁₁₀-(24-Y496) was preceded by the sequence GSPGISGGGGGILDSMGR (SEQ ID No 20).

Production of antibodies that recognise the M_{110} and M_{21} subunits of chicken gizzard $PP1_{M}$.

Polyclonal sheep antibodies to the PP1_M holoenzyme were raised in the Scottish Antibody Production Unit (Carluke, Ayrshire, U.K.). Antibodies which recognise the M₁₁₀ subunit specifically were obtained by passing the antiserum down a 4 ml affinity column comprising 40 mg of M₁₁₀-(R714-I1004) coupled covalently to 1g of dried CNBr-activated Sepharose 4B (Sigma). After washing with 10 column volumes of 50 mM Tris/HCl pH 7.5, 1% (by mass) Triton X-100, 0.1 mM EGTA, 0.1% (by vol) 2-mercaptoethanol (Buffer B) plus 0.5 M NaCl, followed by 10 volumes of Buffer B plus 1 M LiBr, the anti-M₁₁₀ antibody was eluted with 50 mM glycine pH 2.0, neutralised immediately with 1 M Tris/HCl pH 8.0 and stored in aliquots at -80°C. Antibodies which recognise the M₂₁ subunit specifically were obtained in an

identical manner, except that the affinity column comprised about 40 mg of the expressed chicken gizzard M_{21} subunit coupled to 6 g (dry weight) of CNBr-activated Sepharose.

5 Removal of the M_{21} subunit from $PP1_{M}$.

PP1_M (0.01 ml, 0.4U/ml) was dissociated by incubation for 30 minutes with 500 μ M arachidonic acid [25] and then for 30 minutes with 0.08 ml of packed Protein G-Sepharose coupled to 0.08 mg of affinity purified anti-M₂₁ antibody. The Protein G-Sepharose was pelleted, and the supernatant diluted at least 15-fold to allow the M₁₁₀ subunit and PP1c to recombine. The M₁₁₀-PP1c complex was further purified by gel filtration on Superose 12 (30 x 1 cm) to ensure complete removal of any free PP1c.

RESULTS.

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Identification of a PP1c-interaction domain on the G_{M} -subunit of PP1 $_{GM}$. The amino acid sequence of rat hepatic G_L is 23% identical (39% similar) to residues 1-286 of G_M from human skeletal muscle [7]. There is no homology over the first 63 residues but identity is >40% over the regions 63-86, 144-166 and 186-227 of G_M suggesting that one or more of these sequences comprise a PP1-binding domain. Fusion proteins in which GST was linked to fragments of G_M were therefore tested for their ability to bind to PP1c. GST-G_M-(E2-D118) (Fig 1) and GST-G_M-(E2-P243) (data not shown), but not GST-G_M- (H100-P350) or GST itself (Fig 1) interacted with PP1 in Far Western experiments, indicating that the first 118 residues of G_M contain a PP1c-binding domain. Moreover, a proteolytic fragment derived from GST-G_M-(E2-D118) whose molecular mass was 5 kDa less than GST-G_M-(E2-D118), but not a proteolytic fragment that was 6 kDa smaller, also interacted with PP1c (Fig 1). Taken together, the observations suggested that the region comprising residues 63-86 was likely to bind to PP1c. We therefore synthesised $G_{M^-}(G63-T93)$ and examined its effect on the enzymatic properties of $PP1_{GL}$, the form of PP1 associated with rat hepatic protein-glycogen particles.

- The interaction of PP1c with G_L suppresses the dephosphorylation of muscle 5 glycogen phosphorylase by 80% and enhances the dephosphorylation of muscle glycogen synthase by 2-3 fold [21, 26]. Disruption of the characteristic properties of hepatic PP1_{GL} can therefore be monitored very simply by changes G_M-(G63-T93) induced a sixfold increase in the in its specificity. phosphorylase phosphatase activity of PP1_{GL}, the concentrations required for 10 50% activation being 30 nM (Fig 2). G_M-(G63-T93) also prevented bacterially expressed GST-G_L from suppressing the phosphorylase phosphatase activity of PP1c (data not shown). However, G_M-(G63-T93) had no effect on the glycogen synthase phosphatase activity of PP1_{GL}, nor was there any alteration of the other characteristic properties of PP1_{GL}, namely allosteric inhibition of the 15 glycogen synthase phosphatase activity by phosphorylase a and binding to glycogen (data not shown). Thus the interaction of G_{M} -(G63-T93) with PP1_{GL} does not displace G_L from PP1c.
- G_M-(G63-T93) also increased the phosphorylase phosphatase activity of PP1c, indicating that it binds to PP1c, rather than to G_L. However, the maximal stimulation was only 37 + 1.4% (SEM for three experiments), establishing that far greater activation of PP1_{GL} is explained by the ability of G_M-(G63-T93) to overcome the suppressive effect of G_L on the phosphorylase phosphatase activity of PP1c. Several other peptides, including a 32 residue peptide related to the C-terminus of ribosomal protein S6([G245,G246]S6[218-249]), G_M-(S40-Y55) and G_M-(E81-T93) (data not shown), had no effect on the phosphorylase phosphatase activity of PP1_{GL} or PP1c at concentrations up to 10 μM.

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The peptides G_{M} -(G63-K80) and G_{M} -(G63-N75) also increased the phosphorylase phosphatase activity of PP1_{GL}, but were less effective than G_{M} -(G63-T93) and higher concentrations were needed (Fig 2). G_{M} -(G63-K80) and G_{M} -(G63-N75) did not increase the phosphorylase phosphatase activity of PP1c significantly at concentrations up to 10 μ M (data not shown). The phosphorylation of G_{M} at Ser67 by cyclic AMP-dependent protein kinase (PKA) triggers the dissociation of PP1 from G_{M} in vitro and in vivo [18] and phosphorylation of the peptide G_{M} -(G63-N75) at Ser67 prevented it from increasing the phosphorylase phosphatase activity of PP1_{GL} (Fig 2A). The increase in phosphorylase phosphatase activity observed at the highest phosphopeptide concentrations (10 μ M) may be explained by trace contamination (<10%) with dephosphopeptide, resulting either from incomplete phosphorylation of Ser67 or slight dephosphorylation during the assay.

15 Identification of a PPI-interaction domain on the M_{110} subunit.

Antibodies were prepared that recognised either the M_{110} or M_{21} subunits of the myosin-associated form of PP1 (PP1_M) from chicken gizzard (Fig 3A). Removal of the M_{21} subunit using the M_{21} -specific antibody (Fig 3B and see Methods) did not affect the activity of PP1_M towards MLC₂₀ or phosphorylase, the MLC₂₀ phosphatase:phosphorylase phosphatase activity ratio (0.95 \pm 0.03) remaining 15-fold higher than PP1c (Fig 3B). The M_{21} subunit bound to M_{110} , but had no effect on the MLC₂₀ phosphatase or phosphorylase phosphatase activity of PP1c and did not bind to PP1c (D. Johnson unpublished). Thus M_{110} is solely responsible for enhancing the dephosphorylation of MLC₂₀ and suppressing the dephosphorylation of glycogen phosphorylase by PP1c [9].

In order to identify which region(s) of M_{110} modulates the specificity of PP1c, fusion proteins were constructed consisting of glutathione S-transferase (GST) followed by fragments of the M_{110} subunit. After expression in E. coli and

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purification by affinity chromatography on glutathione-Sepharose, the fusion proteins were cleaved with thrombin to release GST from fragments of the M_{110} subunit (Fig 4 and see Methods). M_{110} -(M1-E309), which contains seven 33 residue ankyrin repeats located between residues 39-296, modified the specificity of PP1c in a similar manner to M_{110} itself, increasing activity towards MLC_{20} about 3-fold (Fig 5A) and suppressing activity towards glycogen phosphorylase by about 80% (Fig 5B). The concentration of M_{110} -(M1-E309) required to activate the MLC_{20} phosphatase activity maximally (0.1 nM) was similar to the PP1c concentration in the assay, indicating an extremely high affinity for PP1c. M_{110} -(M1-A150) modified the specificity of PP1 similarly, but 10-fold higher concentrations were needed compared to M_{110} -(M1-E309) (Figs 5C and 5D).

If the GST tags were not cleaved with thrombin, a 10-fold higher concentration of M_{110} -(M1-E309) was needed to modulate the substrate specificity of PP1c, while M_{110} -(M1-A150) was unable to stimulate the MLC₂₀ phosphatase activity of PP1c at all (data not shown). GST itself did not interact with PP1c (Fig 1), had no effect on either the MLC₂₀ phosphatase or phosphorylase phosphatase activity of PP1c (data not shown), and therefore was not removed from the solution after cleavage of the fusion proteins with thrombin.

In contrast to M_{110} -(M1-E309), M_{110} -(D39-E309) failed to stimulate the MLC_{20} phosphatase activity of PP1c, or to inhibit its phosphorylase phosphatase activity (Figs 5A and 5B), suggesting that the extreme N-terminus of the M_{110} subunit (i.e. before the start of the ankyrin repeats) might be important in modulating the specificity of PP1c. The peptide M_{110} -(M1-F38) was therefore synthesized and found to stimulate the MLC_{20} phosphatase activity of PP1c to the same extent as M_{110} -(M1-E309), although the concentration required for half maximal activation (10 nM) was at least 100-fold higher (Fig 5A). M_{110} -(M1-F38) stimulated the dephosphorylation of heavy meromyosin in a

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similar manner to the dephosphorylation of MLC_{20} (data not shown). However, like M_{110} -(D39-E309), M_{110} -(M1-F38) did not inhibit the phosphorylase phosphatase activity of PP1c (Fig 5B). These observations suggested that residues beyond 38 were needed to suppress phosphorylase phosphatase activity. Consistent with this, M_{110} -(L24-Y496) was less effective than M_{110} -(M1-A150) or M_{110} -(M1-E309) in stimulating the MLC_{20} phosphatase activity of PP1c, but inhibited the phosphorylase phosphatase activity of PP1c in a similar manner to M_{110} -(M1-A150) (Figs 5C and 5D).

Although M₁₁₀-(D39-E309) and M₁₁₀-(M1-F38) had no effect on the phosphorylase phosphatase activity of PP1c when each peptide was included individually in the assays at concentrations up to 1 μM (Fig 5), a 39 ± 2% inhibition (SEM n=4) was observed when both peptides were both present at 1 μM. Surprisingly, M₁₁₀-(D39-E309) prevented (IC50 = 0.1 M) M₁₁₀-(M1-F38) from stimulating the MLC₂₀ phosphatase activity of PP1c (data not shown). Thus M₁₁₀-(D39-E309) plus M₁₁₀-(M1-F38) do not faithfully mimic the effect of M₁₁₀-(M1-E309).

We have reported previously that the M_{110}/M_{21} complex suppresses the dephosphorylation of glycogen synthase by PP1c [9] and, consistent with this finding, the dephosphorylation of glycogen synthase was also inhibited by M_{110} -(M1-E309) (Fig 6B). However, the dephosphorylation of glycogen synthase was greatly enhanced by M_{110} -(M1-F38) (Fig 6A).

25 The binding of G_M and the M_{110} subunit to PPIc is mutually exclusive.

In order to investigate whether G_M binds to the same region of PP1c as M_{110} , we next examined the effect of G_M -(G63-T93) on the properties of PP1_M. G_M -(G63-T93) at 10 μ M increased the phosphorylase phosphatase activity of PP1_M by about 7-fold and suppressed its MLC₂₀ phosphatase activity by 60-65%

(Fig 7A), indicating that the distinctive properties of PP1_M had been disrupted. Gel-filtration experiments confirmed that $G_{M^-}(G63-T93)$ had displaced the M_{110} subunit from PP1_M, dissociating it to PP1c (Figs 7B and 7C). $G_{M^-}(G63-T93)$ also prevented $M_{110^-}(M1-F38)$ or $M_{110^-}(M1-E309)$ from stimulating the MLC₂₀ phosphatase activity of PP1c (Fig 8A), and prevented $M_{110^-}(M1-E309)$ from suppressing the phosphorylase phosphatase activity of PP1c (Fig 8B).

Conversely, the presence of $10 \mu M M_{110}$ -(M1-F38) increased the phosphorylase phosphatase activity of PP1_{GL} by 3.5- fold. This resulted from the partial dissociation to PP1c, because the enhanced phosphorylase phosphatase activity was not associated with glycogen, but recovered in the supernatant after centrifugation of the glycogen-protein particles (not shown).

DISCUSSION.

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We have identified a region on G_M that binds to PP1c (Fig 9). The peptides G_M -(G63-T93), G_M -(G63-K80) and G_M -(G63-N75) all prevented G_L from suppressing the dephosphorylation of glycogen phosphorylase by PP1c and two lines of evidence indicate that these peptides interact with PP1c and not with G_L .

Firstly, the PP1c-catalysed dephosphorylation of glycogen phosphorylase is stimulated slightly by G_{M} -(G63-T93).

Secondly, PP1c crystallises in the presence of G_M-(G63-K80) or G_M-(G63-N75) in a different form than is observed in the absence of these peptides. PKA phosphorylates G_M at Ser67 and the introduction of a negative charge directly into the PP1c-binding domain explains why phosphorylation of Ser67 triggers the dissociation of G_M from PP1c [18]. Phosphorylation of G_M-(G63-N75) at

Ser67 also prevented this peptide from interacting with PP1 in the PP1_{GL}

complex (Fig 2).

Although G_{M} (G63-T93) prevented G_{L} from suppressing the dephosphorylation of glycogen phosphorylase by PP1c, it did not dissociate G_L from PP1c, nor did it affect the other characteristic properties of PP1_{GL}. Moreover, unlike G_L, G_M-(G63-T93) did not itself suppress the phosphorylase phosphatase activity of PP1c, but actually enhanced it slightly. These observations demonstrate that another region(s) on G_L must interact with PP1c and that this other region(s) may play an important role in modulating the substrate specificity of PP1c. The presence of a second PP1c binding site in G_M/G_L would be somewhat 10 analogous to the situation found in inhibitor-1 and DARPP which also contain two PP1-binding sites, high (nM) affinity binding being generated by the conjugation of two low affinity binding sites that, individually, only interact with PP1 at μ M concentrations [28]. The second PP1c-binding site on G_M/G_L might correspond to one of the other regions where G_M and G_L show >40%15 identity (residues 144-166 and 186-227 of human G_M). G_M-(H100-P350) was not recognised by PP1c in Far Western experiments (Fig 1) this result is not definitive because G_M-(H100-P350) may only interact with PP1c weakly. Alternatively, G_M-(H100-P350) might not fold correctly or fail to renature after SDS/polyacrylamide gel electrophoresis. 20

However, it is also possible that residues 144-166 and 186-227 of G_M do not represent part of the second PP1c-binding domain, but part of the glycogen-binding domain. In this connection it should be recalled that residues 144-166 and 186-227 are the regions showing greatest similarity (25% identity) to GAC1, which appears to be a homologue of G_M/G_L in budding yeast [7, 27, 28]. Curiously, GAC1 does not contain a region homologous to residues 63-93 of G_M/G_L . It would clearly be of interest to compare the effect of GAC1 on the enzymatic properties of PP1c with those of G_M and G_L .

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We have also identified a region on the M_{110} subunit that binds to PP1c. An N-terminal fragment, M100-(M1-E309), enhanced the PP1c-catalysed dephosphorylation of MLC₂₀ and suppressed the dephosphorylation of glycogen phosphorylase in a similar manner to M_{110} itself (Fig 5). However, unlike M_{110} , this fragment does not bind to myosin. Thus the region which enhances the dephosphorylation of MLC₂₀ is distinct from the myosin-binding domain.

The fragment M₁₁₀-(M1-E309) contains seven ankyrin repeats lying between residues 39 and 296. However, M₁₁₀-(D39-E309) was ineffective as an activator of the MLC₂₀ phosphatase activity of PP1c or as an inhibitor of the phosphorylase phosphatase activity, and this led to the finding that a peptide comprising the N-terminal 38 residues of the M₁₁₀ subunit enhances the dephosphorylation of MLC₂₀ to the same extent as M_{110} -(M1-E309), although with lower potency. However, M_{110} -(M1-F38) did not inhibit the dephosphorylation of glycogen phosphorylase by PP1c suggesting that residues beyond 38 are required to suppress this activity. This view was reinforced by the finding that, although neither M₁₁₀-(M1-F38) nor M₁₁₀-(D39-E309) inhibited the phosphorylase phosphatase activity of PP1c when present individually, inhibition was observed in the presence of both peptides. Moreover M_{110} -(D39-E309) actually prevented M_{110} -(M1-F38) from stimulating the dephosphorylation of MLC₂₀.

These observations suggest that M_{110} -(D39-E309) can bind to M_{110} -(M1-F38) and/or PP1c. An interaction with PP1c seems likely because it has been found that M_{110} -(D39-E309) can enhance the phosphorylase activity of PP1_{GL}. The presence of a second PP1-binding site in the ankyrin-repeat domain of the M_{110} subunit is also supported by the observation that higher concentrations of M_{110} -(M1-A150) and M_{110} -(M1-E309) are needed to inhibit the phosphorylase phosphatase activity of PP1c than are required to stimulate its MLC_{20} phosphatase activity (see Fig 5). The presence of at least two PP1-binding sites

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may explain why the M_{110} subunit and PP1c interact at picomolar concentrations. The ankyrin repeat domain might suppress the dephosphorylation of some substrates (such as glycogen phosphorylase) by a steric mechanism, preventing them from gaining easy access to the catalytic centre. This scenario could explain why the dephosphorylation of glycogen synthase is greatly enhanced by M_{110} -(M1-F38) yet suppressed by M_{110} -(M1-E309) (Fig 6).

 G_{M} -(G63-T93) abolished the distinctive properties of PP1_M (Fig 7A), prevented M_{110} -(M1-F38) or M_{110} -(M1-E309) from modulating the substrate specificity of PP1c (Fig 8) and displaced the M_{110} subunit from PP1_M (Fig 7B). In addition, the peptide M_{110} -(M1-F38), was capable of displacing G_L from PP1_{GL}. These findings indicate that the binding site(s) on PP1c for G_M and the M_{110} subunit are likely to overlap, explaining why different forms of PP1 contain a single PP1-targeting subunit. The three-dimensional structure of PP1c isoforms have recently been solved to high resolution [29,30], and PP1c crystallises in different forms in the presence of G_M -(G63-N75) or G_M -(G63-K80) or M_{110} -(M1-F38) than in the absence of these peptides.

Consistent with the results presented here, Gailly et al [31] have recently shown that M₁₁₀-(M1-F38) or M₁₁₀(M1-E309) enhance the ability of PP1c to stimulate the relaxation of microcystin-contracted permeabilised portal vein, while G_M-(G63-T93) inhibits the ability of PP1_M to induce the relaxation of this smooth muscle. G_M-(G63-T93) also slowed the relaxation of permeabilised femoral artery, indicating that it competes with the endogenous M₁₁₀ subunit for PP1c [31]. Thus the PP1c-binding peptides described constitute useful pharmacological agents with which to explore the role and regulate the activity of PP1 in cell regulation.

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Example 2: Structural basis for the recognition of regulatory subunits by the catalytic subunit of protein phosphatase 1

MATERIALS AND METHODS

Crystallisation and Data Collection

The catalytic subunit of PP1 1 was overproduced in Escherischia coli and purified as described previously (Alessi et al., 1993; Barford and Keller, 1994). The $G_M[G63-N75]$ peptide, variants of this peptide in which Val 66' or Phe 68' were changed to Phe, and the peptides $M_{110}[1-38]$ and $M_{110}[1-35]$ were synthesised on an Applied Biosystems 430A peptide synthesiser and purified by chromatography on a C18 column (Johnson et al., 1996) by Mr F.B. Caudwell at Dundee. A three-fold molar excess of G_M[G63-N75] was added to the protein solution (8 mg/ml), which had been previously dialysed against 10 mM Tris-HC1 (pH 7.8), 0.3 M NaCl, 0.4 mM MnCl, and 2 mM DTT. The complex was crystallised at 20°C using the hanging drop vapour diffusion method, by mixing 2 ml of the protein-peptide solution and 2 ml of the precipitant solution containing 2.0 M ammonium sulphate, 2% (w/v) polyethylene glycol 400,100 mM HEPES (pH 7.5) and 2 mM DTT. These conditions are very much in contrast to the relatively low ionic strength conditions from which the monoclinic PP1c crystals grew (Barford and Keller, 1994; Egloff et al., 1995). Crystals appeared after 3 months as a cluster. Individual crystals removed from the cluster had dimensions of $\sim 25 \mu m \times 25$ μ m x 5 μ m. Crystals were frozen in a 100 K nitrogen gas stream and stored. Prior to freezing, crystals were incubated in a cryoprotectant solution consisting of an equilibration buffer; 2.0 M ammonium sulphate, 2% (w/v) PEG 400, 100 mM HEPES (pH 7.5) with increasing amounts of glycerol in steps of 7%, 15%, 22% and 30% (v/v).

A partial data set to 3.0 Å was collected on Beam Line PX 9.6, SRS,

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Daresbury, using a 30 cm diameter Mar Research image plate system. Data were processed and scaled using DENZO and SCALEPACK (Otwinowski, 1993). The crystal system is tetragonal with point group symmetry P422 and unit cell dimensions a = b = 62.50 Å, c = 361.30 Å. Systematic absences indicate a 21 screw axis along b. The Matthews coefficient was 2.38 Å₃ per Dalton, assuming 2 molecules per asymmetric unit. A second data-set was collected on BL4 at the ESRF, Grenoble. Substantial radiation damage was observed during data collection requiring that three crystals were used in total. Data collected from four crystal at Daresbury and the ESRF were merged together in SCALEPACK. Details of the data collection and processing statistics are given in Table 1.

Structure determination

The structure of the PP1- $G_M[63-75]$ complex was solved by molecular replacement using as a model the protein atoms coordinates of the 2.5Å refined structure of the catalytic subunit of PP1 γ 1 determined by MAD methods (Egloff *et al.*, 1995). Rotation and translation functions searches were performed with AMORE (Navaza, 1992). Using data between 8 and 3 Å resolution, the peak in the rotation search was 6.7 standard deviations (SD) above the mean. The translation search was best performed using data between 8 and 3.5 Å, giving a maximal peak at 13.8 SD above the mean for the space group P41212. After the first rigid body refinement performed in AMORE, the R-factor was 0.494 and the correlation factor 0.30.

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Crystallographic Refinement

The solution from molecular replacement was optimized by 20 cycles of rigid body refinement performed with X-PLOR version 3.1 (Brunger, 1992), using data between 8.0 Å and 3.0 Å resolution. After a round of conjugate gradient

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positional refinement and simulated annealing molecular dynamics to 2000 K, followed by 25 cycles of grouped B-factor refinement (2 B-factor groups for each residue), the R factor (respectively free-R) was 0.295 (0.367). Fourier difference maps (Fo-Fc) and (3Fo-2Fc) revealed the presence of three strong peaks at (over three-times the sigma level of the map) at the catalytic site of PP1c. From the previously refined PP1c-structure, we identified two as manganese and iron ions. The third one, occupying the position of the tungstate ion in the PP1c-WO4 complex, was identified as sulphate. The initial difference Fourier maps also revealed strong electron density near the N-terminus of β 14. The maps were improved by applying non-crystallographic symmetry 2-fold averaging using PHASES (Furey and Swaminathan, 1990). As shown in Fig. 1A, residues Val 66', Ser 67' and Phe 68' of the $G_M[63-75]$ peptide were identified in the averaged map. These 3 residues, as well as the 2 metal and sulphate ions were built in each molecule, using the program TURBO-FRODO (Roussel and Cambillau, 1992). Refinement of this structure was performed by repeated rounds of manual rebuilding followed by conjugate gradient positional refinement and grouped B-factor refinement using X-PLOR. The final model contains protein residues Lys 6 to Ala 299 and peptide residues Arg 65' to Ala 69' in molecule 1, and protein residues Asn 8 to Lys 297 and peptide residues Gly 63' to Ala 69' in molecule 2. A few well defined water molecules were also observed in both initial (3Fo-2Fc) and (Fo-Fc) electron density maps. Eventually, 14 water molecules that were above 3 sigma in the (Fo-Fc) difference map, within hydrogen bond of the PP1-peptide complex or another solvent molecule and present in both molecules, were included in the model. The crystallographic and refinement data are summarized in Table 1. Representative electron density from the peptide before and after refinement is shown in Figure 10A and 10B, respectively. Solvent accessible surface areas were calculated using the method of Lee and Richards (1974).

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Purification and assay of PP1.

PP1c was isolated from the rabbit skeletal muscle PP1-G_M complex as described previously (Johnson *et al*, 1996). Glycogen particles isolated from rat liver (Schelling *et al*, 1988) served as the source of PP1-G_L. The dephosphorylation of glycogen phosphorylase (10 μM) and the isolated MLC₂₀ of smooth muscle myosin (l μM) by PP1c was carried out as described previously (Cohen *et al.*, 1988; Alessi *et al.*, 1992).

10 Table 1. Crystallographic data and refinement statistics

Crystallographic data:				
Space group	P4 ₁ 2 ₁ 2			
Unit cell parameters (Å)	a = b = 62.50; c = 361.30			
Number of molecules per asymmetric unit	2			
Temperature (K)	100			
Total measured reflections	290671			
Number of unique reflections	15509			
Mean I/s(I)	7.5			
Completeness (%)	87			
Overall R-merge (%)	14.7			
Refinement statistics:				
Number of reflections used for refinement	13078			
Resolution range (Å)	8.0-3.0			
R-work	0.223			
R-free	0.308			
Number of residues	protein peptide			

Molecule 1	294 (Lys 6 to 6 (ARVSFA) (SEQ ID No 21) Ala 299) 6 (RRVSFA) SEQ ID No 3)
Molecule 2	290 (Asn 8 to Lys 297)
R.m.s.d. from ideal bond lengths (Å)	0.012
R.m.s.d. from ideal angles (°)	1.863
Number of water molecules	
Molecule 1	7
Molecule 2	7

10 Table 2. PP1-peptide polar interactions

	Peptide atom	Protein atom	Water molecule	Distance (Å)
Molecule 1	Arg 65' O	-	7W	3.2
	Val 66' N	Asp 242OD2 (**)		3.0
	Ser 67' N	Leu 289 O		3.3
	Ser 67' OG		7W	2.7
	Ser 67' O	Cys 291 N (*)		3.2
	Ala 69' N	Cys 291 O (*)		2.8
Molecule 2	Arg 64' NH1	Glu 287 O		2.6
		(**)		
	Arg 65' O		7W	2.8
	Val 66' N	Asp 242 OD2 (**)		3.2
	Ser 67' N	Leu 289 O (*)		3.1
	Ser 67' OG		7W	2.6
	Ser 67'	Cys 291 N (*)		3.0

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	Ala 69' N	Cys 291 O (*)	3.3			
Table 2. PP1-peptide hydrophobic interactions						
	Peptide residues	Protein residues				
	Val 66'	Ile 169 (*), Leu 243 (*), D242 (**), Leu 289 (*), Cys 291 (*)				
	Phe 68'	Phe 257 (*), Cys 291 (*), Phe 293 (*)				
	Ala 69'	Met290 (**)				

The star (*) indicates residues absolutely conserved in all protein phosphatase 1 sequences available so far, the double start (**) the residues mostly conserved (from sequence alignment from Barton et al, 1994).

RESULTS AND DISCUSSION

Structure Determination.

Crystallographic data to 3.0 Å were measured at the ESRF beam-line BL4 at Grenoble and at PX9.6, Daresbury (Table 1). The relatively high merging R-factors and low I/(I values of the crystallographic data results from the weak diffraction observed from the PP1- $G_M[63-73]$ crystals. This is attributable to both the small crystal size ($\sim 25~\mu m$ by $25~\mu m$ by $5~\mu m$) and long c-axis of the unit cell. In addition, the high x-ray photon dose required to obtain usable diffraction images resulted in x-ray radiation damage to the crystals, despite being maintained at a temperature of 100 K during the course of the experiment. The structure was solve by the molecular replacement method using as a search model the 2.5 Å refined coordinates of PP1c (Egloff et al., 1995). Phases obtained from a single cycle of simulated annealing refinement

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of the protein coordinates alone using X-PLOR Brunger, 1992), and improved by 2-fold non-crystallographic symmetry averaging and solvent flattening, were used to calculate an electron density map. This map revealed clear density corresponding to residues Val 66', Ser 67' and Phe 68' (where 'denotes residues of the peptide) of the G_M peptide and provided a starting point for further refinement of the PP1- G_M peptide complex (Fig. 10A). The final model of the complex was refined at 3.0 Å resolution with a crystallographic R-factor of 0.22 and R-free of 0.31 (Fig. 10B). The two molecules of PP1c within the asymmetric unit are similar with a root mean square deviation between main chain atoms of 0.6 Å. Residues 6 to 299 and 8 to 297 from molecules 1 and 2 respectively, are visible in the electron density map. Similar to the structures of native $PP_{\gamma 1}$ (Egloff et al., 1995) and $PP1\alpha$ in complex with microcystin LR (Goldberg et al., 1995), residues C-terminal to 299 are disordered.

15 Overall Structure of PP1

The conformation of PP1c in the PP1- G_M complex is virtually identical to that of native PP1c in complex with tungstate (Egloff et al., 1995) with a root mean square deviation between equivalent main-chain atoms of 1.0 Å. PP1c is folded into a single elliptical domain consisting of a central β -sandwich of two mixed β -sheets surrounded on one side by 7α -helices and on the other by a sub-domain consisting of 3α -helices and a 3 stranded mixed α -sheet (Fig. 2A, B). The interface of the three β -sheets at the top of the β -sandwich creates a shallow catalytic site channel. Three loops connecting β -strands with α -helices within a β - α - β - α - β motif in sheet 1 (strand order β 4- β 3- β 2- β 13- β 14) together with loops emanating from the opposite β -sheet (sheet 2; strand order, β 1- β 5- β 6- β 10- β 12- β 11) provide the catalytic site residues. The catalytic site of PP1 contains a binuclear metal site consisting of Mn²⁺ and Fe²⁺ (Egloff et al., 1995) and, in the PP1- G_M complex, oxygen atoms of a sulphate ion of crystallisation coordinate both metal ions, similar to that seen in the PP1-tungstate (Egloff et al., 1995) and PP2B-phosphate complexes (Griffith et

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al, 1995).

$PP1c-G_{M}[63-75]$ Peptide Interactions

Six residues of the $G_M[63-75]$ peptide (Arg 64' to Ala 69') are clearly visible in the electron density map of the complex of molecule 2, the remaining residues are not visible and assumed to be disordered (Fig. 10B). Density is not visible for Arg 64' of the peptide bound to molecule 1, otherwise equivalent residues of the peptide are similar within the two complexes. The six residues (RRVSFA) (SEQ ID No 3) of the $G_M[63-75]$ peptide in complex 2 adopt an extended conformation and bind to a hydrophobic channel on the protein surface with dimensions 25 Å by 10 Å that is formed at the interface of the two β -sheets of the β -sandwich opposite to the catalytic site channel and therefore remote from the catalytic site (Fig. 11A). The residues that form this channel occur on three regions of PP1c, namely (i) the N-terminus of 5 and $\beta 5/\beta 6$ loop of sheet 2; (ii) the three edge β -strands of sheet 2: β 10, β 12, β 11 and (iii) β 13, the $\beta 13/\beta 14$ loop and $\beta 14$ of the edge of sheet 1 (Fig. 11A). The total solvent accessible surface area buried on formation of the complex is 980 A2. Three residues of the peptide (Ser 67' to Ala 69') form a β -strand which is incorporated into β -sheet 1 of PP1c as a sixth β -strand parallel to the N-terminus of the edge β -strand, β 14 (residues Leu 289 to Leu 296) (Fig. Main-chain atoms of Ser 67' and Ala 69' form H-bonds to the main-chain atoms of residues of β 14. In addition, the main-chain nitrogen of Val 66' forms a H-bond with the side-chain of Asp 242. interactions include the guanidinium group of Arg 64' with the mainchain carbonyl of Glu 287 and a salt bridge to Asp 166. Both Asp 166 and Asp 242 A water molecule bridges the are invariant in mammalian PP1 genes. main-chain carbonyl of Arg 65' and side-chain hydroxyl of Ser 67' with the main-chain carbonyl of Thr 288 of PP1c (Fig. 11C). A notable feature of the peptide binding site is the presence of a negatively charged region created by seven acidic residues (with one Lys residue) surrounding the hydrophobic channel at the N-terminus of the peptide in the vicinity of Arg 64' and Arg 65' that includes Asp 166 and Asp 242 (Fig. 11D). This would suggest a favourable electrostatic environment for the side chains of Arg 64' and Arg 65'.

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The predominant interactions between the peptide and PP1c involve hydrophobic contacts between the side chains of Val 66' and Phe 68' and solvent exposed, invariant, hydrophobic residues of PP1c that form the hydrophobic channel (Fig. 11C, E). In particular, the binding site for the side chain of Val 66' is formed from the side chains of Ile 169, Leu 243, Leu 289 and Cys 291, whereas that for the side chain of Phe 68' is formed from the side chains of Phe 257, Cys 291 and Phe 293. Details of peptide-PP1c contacts are given in Table 2. The structure of the $G_M[63-75]$ peptide binding site is likely to be conserved in other forms of PP1 from diverse species. Each hydrophobic residue of PP1c that interacts with the Val 66' and Phe 68' residues of the $G_{M}[63-75]$ peptide are invariant and the acidic residues that surround the N-terminus of the peptide binding site are highly conserved amongst all isoforms of PP1 from species as diverse as yeast, Drosophila, mammals and higher plants (Barton et al., 1994). However, since these residues are not conserved within the PP2A and PP2B sequences, these proteins will not recognise PP1-regulatory subunits.

Presence of an (R/K) (V/I) x F Motif in other PP1c Regulatory Proteins

Over a dozen regulatory subunits of PP1c are now known which appear to bind to PP1c in a mutually exclusive manner that suggests either an overlapping binding site or sites. Sequence comparisons of these subunits reveals little similarity except for the motif (R/K) (V/I) x F, that is not only present in $G_M[63-75]$ but also in G_M , G_L , M_{110} , NIPP-1, p53BP2, and an RNA splicing factor (Fig. 12A). Moreover, a 38 residue peptide from the 110kDa M_{110} subunit that binds to PP1c contain this motif (Johnson et al, 1996), as do

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fragments of NIPP-1 (Beullens et al., 1992; Van Eynde et al., 1995), an RNA splicing factor (Hirano et al., 1996) and p53BP2 (Helps et al., 1995). A 32 residue peptide from p53BP2, which contains this motif, disrupted the interaction of the M_{110} subunit with PP1c, as shown by a decrease in the rate of dephosphorylation of the MLC₂₀ subunit of smooth muscle myosin and by an increase in the rate of dephosphorylation of glycogen phosphorylase (Fig. 13A). This peptide also disrupted the interaction of the G_L subunit with PP1c, as shown by an increase in the rate of dephosphorylation of glycogen phosphorylase (Fig. 13B). Peptides comprising the motif (R/K) (V/I) x F are thus encompassed within the scope of the invention.

In further support of the notion of a common PP1c recognition motif present within PP1-binding proteins, previous studies had revealed that the sequence KIQF (SEQ ID No 22) (similar to the R/KVxF motif) at the N-terminus of inhibitor 1 and its homologue DARPP-32 (Fig. 12A) is necessary for mediating the inhibitory effects of these proteins. Loss of Ile 10 of the KIQF (SEQ ID No 22) motif of inhibitor 1 disrupts the inhibitory effects on PP1c by phospho-inhibitor-1 (Aitken and Cohen, 1984; Endo et al., 1996) and the binding of either dephospho-inhibitor-1 or phospho-inhibitor-1 to PP1c (Endo et al., 1996). A similar result was found on disrupting the equivalent residue (Ile 9) of DARPP-32 (Hemmings et al., 1990; Desdouits et al., 1995). These results were interpreted to indicate that inhibitor-1 and DARPP-32 bind to PP1 through two low affinity binding sites, one that encompasses the sequence KIOF (SEO ID No 22) and another which includes the phosphorylated Thr residue (35 in I-1, 34 in DARPP-32) and which presumably binds at the catalytic site. Analysis of the PP1-G_M[63-75] complex structure suggests that an isoleucine residue could be readily accommodated within the peptide binding site in place of Val 66' such that the additional methyl group on Ile compared to Val would contribute to favourable van der Waals interactions between the peptide and Leu 243 and Cys 291 of PP1. More bulky hydrophobic residues

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such as Leu, Met and Phe cannot be accommodated, however. It is interesting to note that as well as the (R/K) (V/I) x F motif shared by PP1-regulatory subunits, the four residues N-terminal to this motif contain an abundance of basic residues. These residues may provide further favourable interactions with the negative electrostatic surface potential at the N-terminus of the $G_M(63-75)$ peptide binding site of PP1c (Fig. 11D).

Mutagenesis of the R/K) $(V/I) \times F$ motif

The structural studies presented here suggest a dominant role for Val 66' and Phe 68' in stabilising the interaction between $G_M[63-75]$ and PP1c and this notion is further reinforced by the finding that other PP1-regulatory subunit sequences contain an (R/K)(V/I) x F motif yet share little overall sequence similarity. To test the hypothesis that Val 66' and Phe 68' are required for the interaction of $G_M[63-75]$ with PP1c and also that the KVKF (SEQ ID No 5) sequence present within the $M_{110}[M1-F38)$ peptide is important in mediating its interaction with PP1c, we synthesised variations of the G_M and M_{110} peptides where the R/KVxF motif was disrupted. The two variants of the G_M peptide were Val 66' and Phe 68' to Ala substitutions. In order to disrupt the (R/K)(V/I) x F present within the M_{110} peptide, a peptide corresponding to residues Met 1 to Lys 35 was synthesised which no longer contains the sequence VKF of the VxF motif, which is present at residues 36-38.

The results for the $M_{110}[1-38]$ and $M_{110}[1-35]$ peptides (Figs. 14, 15) are unequivocal. Whereas $M_{110}[1-38]$ stimulates the myosin light chain phosphatase activity of PP1c with a half-maximal effect at 10 nM reaching maximal (3-fold) activation at a peptide concentration of 1 μ M as reported previously (Johnson et al, 1996), the $M_{110}[1-35]$ peptide was at least 104-fold less effective at activating PP1c (Fig. 14). Unlike $M_{110}[1-38]$, the $M_{110}[1-35]$ peptide was also unable to activate the phosphorylase phosphatase activity of liver PP1- G_L . This latter result suggests two conclusions. Firstly, that although $M_{110}[1-38]$ is able

to bind to PP1c and disrupt the interactions between PP1c and the G_L -subunit, hence reversing the inhibitory effects of G_L on the ability of PP1c to dephosphorylate phosphorylase, loss of the VKF sequence in the $M_{110}[1-38]$ peptide abolishes the ability of the peptide to disrupt this interaction. Secondly, the recognition site on PP1c for the VKF sequence of the $M_{110}[1-38]$ peptide must overlap with the binding site for the G_L subunit, suggesting that the VKF sequence binds to the same site as the VSF sequence of G_L that is identical with that present in the $G_M[63-75]$ peptide. Similar conclusions may be reached from the results obtained from disrupting the VxF motif within the $G_M[63-75]$ peptide (Fig. 16B). Substitution of Phe 68' for Ala abolishes completely the ability of $G_M[63-75]$ to disrupt the PP1- G_L complex, whereas replacement of Val 66' with Ala reduced the effectiveness of the disruption 100-fold.

Thus preferred peptides may comprise analogues of G_M with substitutions at Val 66' and Phe 68' to some other amino acid such as Ala, so that binding of the PP1c to G_M does not occur and the PP1c is not suitably directed or controlled. Alternatively, suitable peptides could comprise peptides suitable to compete for the binding site(s) of Val 66' and Phe 68' on PP1c. Such peptides can be added in sufficient quantities to compete for the Phe 68' and Val 66' binding site(s) on the PP1c, thereby disrupting the interaction of PP1c and natural G_M . Such peptides could comprise structural analogues of G_M with Phe 68' and Val 66' in the same positions as G_M . Alternatively, other amino acids capable of mimicking the binding of Phe 68' and Val 66' could be used in these locations.

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Regulation of the PP1- G_M Complex by Phosphorylation of Ser 67'

Phosphorylation of Ser 67', corresponding to x of the VxF motif, by PKA promotes dissociation of both G_M and G_M [63-75] from PP1c. In vivo, this provides a mechanism of inhibiting PP1c during stimulation of skeletal muscle

by adrenalin (Dent et al., 1990). The sequence of G_M surrounding Ser 67' (RRVSFA) (SEQ ID No 3) conforms to a consensus PKA recognition sequence. Interestingly, the conformation of the peptide is similar to that of residues 18 to 23 corresponding to the pseudo-substrate sequence of PKI bound to the catalytic site of PKA (Knighton et al., 1990). Although the side chain of Ser 67' is exposed within the PP1c-peptide complex, overall the G_M peptide is buried, and it is unlikely that Ser 67' would be a substrate for PKA when the peptide is bound to PP1c. This would suggest that PKA phosphorylates Ser 67' when G_M is not associated with PP1c and that this phosphorylation prevents the re-association of PP1c with G_M. Since phosphorylation of Ser 67' promotes the dissociation of the PP1-G_M complex both in vivo and in vitro, it is most likely that PKA phosphorylates Ser 67' of G_M by competing with PP1c for the RRVSFA (SEQ ID No 3) sequence. This is consistent with a notion that the PP1-G_M complex exists in dynamic equilibrium with free PP1c and G_M subunits and that phosphorylation occurs on the regulatory subunit during transient dissociation from PP1c. In the PP1c-peptide complex, thè side-chain of Ser 67' adopts the most favourable rotamer conformation. Analysis of the PP1c peptide complex structure suggests that incorporation of a phosphate group onto the side chain of Ser 67' with the same side-chain rotomer conformation would cause steric hindrance between the peptide and Met 290 of PP1 and also introduce a phosphate group into a region of negative charge at the PP1c surface (Fig. 11C, D). This may explain how phosphorylation of Ser 67' prevents peptide association with PP1c, although it should be noted that rotation of the side-chain of Ser 67' would relieve this steric clash.

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A similar mechanism of control may also operate for other PP1-regulatory subunits. For example, NIPP-1 a nuclear inhibitor of PP1, inhibits PP1 with an inhibitory constant of 1 pM (Beullens et al., 1992). Phosphorylation of NIPP-1 by PKA and/or casein kinase 2 in vitro abolishes this inhibition (Beullens et al., 1993; Van Eynde et al., 1994). Although the sites of

phosphorylation on NIPP-1 that mediate these effects are not yet fully characterised it is known that these sites occur within the central ~120 residues of NIPP-1 that incorporates the (R/K)(V/I) x F motif (Van Eynde et al., 1995). Interestingly, a consensus phosphorylation site for PKA (RKNS) (SEQ ID No 23) occurs immediately N-terminal to this motif whereas one casein kinase 2 consensus phosphorylation site occurs between the Val and Phe of the motif and another occurs immediately C-terminal to the Phe residue (TFSEDDE) (SEQ ID No 24) (Van Eynde et al., 1995) (Fig. 12A). It is possible that PKA, casein kinase II or other kinases with similar specificity, release PP1c from inhibition by NIPP-1 by phosphorylating NIPP-1 at sites that block its interaction with the (R/K)(V/I) x F motif recognition site on PP1c.

Model of the PPIc-Phospho-Inhibitor 1 Complex

Our model for the interaction of a (R/K)(V/I) x F motif with PP1c, together 15 with previous kinetic data suggesting that the sequence KIQF (SEQ ID No 22) of inhibitor-1 (Aitken and Cohen, 1984; Endo et al., 1996) and DARPP-32 (Hemmings et al., 1990; Desdouits et al, 1995) interacts with PP1c, allowed us to construct a plausible model of a complex of PP1c with phospho-inhibitor 1. The major assumptions of this model were (1) the KIQF (SEQ ID No 22) 20 sequence of inhibitor-1 binds to the same site as RVSF (SEQ ID No 25) of the G_w[63-75] sequence and (2) that the phosphothreonine residue 35 of phospho-inhibitor 1 binds at the phosphate binding site of the PP1c-catalytic site. Secondary structure predictions of inhibitor 1 (Rost and Sander, 1993; Rost, 1996) suggested that residues 9 to 14 and 23 to 31 adopt β -strand and 25 α-helical conformations, respectively. The prediction of the sequence KIQF (SEO ID No 22) as a β -strand is consistent with our assumption that this region of inhibitor-1 adopts the same conformation as RVSF (SEQ ID No 25) of the G_M peptide when bound to the VxF recognition site of PP1c. We have positioned the residues RRPpTP (SEQ ID No 26) encompassing the pThr 35 30

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site within the catalytic site channel in an extended conformation, with the phosphate group of the pThr 35 occupying the phosphate binding site of the catalytic site and the Oy-atom of Thr 35 equivalent to the solvent exposed oxygen of a dianion that forms a H-bond to the side-chain of the putative general acid His 125 (Egloff et al., 1995; Griffith et al., 1995). The four consecutive Arg residues N-terminal to pThr 35 interact with Asp and Glu residues within an acidic groove of PP1c formed from the $\beta7/\beta8$ loop on one side and the $\beta 10/\beta 11$ loop and $\beta 11$ strand on the other, similar to that proposed by Goldberg et al., (1995) for their model of DARPP-32 bound to PP1c. We propose that residues 20 to 30 of inhibitor-1 form an amphipathic helix which folds around the edge of the β -sandwich of PP1c. The N-terminus of this helix is disrupted by prolines at residues 19 and 23. Pro 19 and Pro 15 are probably responsible for introducing turns into the polypeptide chain that allows the β -strand encompassing the KIQF (SEQ ID No 22) sequence (residues 9 to 14) to connect with the α helix. The model of the phospho-inhibitor 1-PP1c complex is shown in Fig. 16.

Prediction of PP1 Recognition Motifs in Yeast PP1-Binding Proteins

The residues in mammalian PP1c that interact with the sequence RRVSFA (SEQ ID No 3) are conserved in S. cerevisiae PP1 suggesting that the proteins in S. cerevisiae known to interact with PP1 (reviewed by Stark, 1996) probably bind to a similar hydrophobic groove on the surface of the enzyme. Examination of their amino acid sequences revealed that a number of PP1-binding proteins in S. cerevisiae contained putative PP1-binding motifs that were similar to those present in mammalian PP1-binding proteins (Fig. 12A, B). The S. cerevisiae PP1-binding proteins not only contain a V/I x F motif, but also a basic residue equivalent to Arg 64' of G_MV the residue that contacts Asp 166, Leu 289 and the main-chain carbonyl of Glu 287 of PP1c. Several of the S. cerevisiae proteins also contain a further basic residue (His or Lys) at the position equivalent to Arg 65' of G_M . Another striking feature of the

putative PP1-binding sequences in S. cerevisiae is the presence of a basic amino acid between the Val/Ile and Phe residues, as is also found in two mammalian PP1-regulatory subunits, the M_{110} subunit and the p53BP2 (Fig. 12A).

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The S. cerevisiae proteins GAC1 and PIG2 show some homology to residues 140-230 of mammalian G_MV and there is genetic and biochemical evidence that they may function to regulate glycogen metabolism in budding yeast (François et al., 1992). GIP2 also shares sequence similarity with residues 140-230 of mammalian G_M, while YIL045W is an open reading frame in the S. cerevisiae genome whose predicted amino acid sequence shows 41 % sequence identity to GIP2. YIL045W contains two putative PP1-binding motifs and site directed mutagenesis will be needed to establish which (if either) of these sequences binds to PP1c. REG1 and REG2 are PP1-binding proteins that play a role in cell growth and, in the case of REG1, glucose repression (Tu and Carlson, 1995; Tu et al., 1996; Frederick and Tatchell, 1996). GIP1, which also contains two putative PP1-binding motifs, is expressed specifically during meiosis, affects the transcription of late meiotic genes and is essential for sporulation (Tu and Carlson, 1996). SCD5 is a PP1-interacting protein (Tu et al., 1996) that was first isolated as a multicopy suppressor of the inviability of clathrin heavy chain-deficient yeast (Nelson et al., 1996).

The findings herein demonstrate that the short peptide sequence, the (R/K)(V/I)XF motif, is critical for PP1c to interact with its regulatory subunits. PP1c (when complexed to its targeting subunits) plays key roles in the control of many cellular processed and it is reasonable to predict that over 100 pp1-binding proteins may exist in mammalian cells. Protein sequence data-base searching has revealed that the (R/K)(V/I)XF motifs are found in 10% of proteins. Thus if ~ 100 PP1-binding proteins occur in mammalian cells, only 1% of proteins with the (R/K)(V/I)XF motif will be PP1-binding proteins. The

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reasons why only a few proteins with the (R/K)(V/I)XF motif bind to PP1 are numerous. For example, not every residue may be tolerated at position X or immediately N-terminal or C-terminal to this motif. This study has shown that phosphoserine is not tolerated at position X and it is therefore likely that Asp or Glu will not be tolerated either. The structure of the PP1- G_M [63-75] complex suggests that large hydrophobic residues will also be excluded from position X. Moreover, the Val (or Ile) and Phe residues in many (R/K)(V/I)XF motifs will be buried in the hydrophobic core of the protein and hence be unable to interact with PP1, since this motif is predicted to form an amphipathic β -strand conformation. Thirdly, many of the (R/K)(V/I)XF motifs will be in extracellular proteins or extracelluar domains of transmembrane proteins and hence be unable to bind to PP1. Particular feature so the tertiary structure of PP1-binding proteins may allow exposure of this motif on the surface to allow interaction with PP1. Finally, there is evidence that a second PP1-binding site exists on the G_M and M_{110} subunits (Johnson et al., 1996) and the high affinity interaction of PP1c with protein inhibitor-1 is generated by the binding of PP1c to two low affinity sites (Desdouits et al., 1995), one of which is the KIQF sequence belonging to the (R/K)(V/I)XF motif.

The question of how regulatory subunits modulate the substrate specificity of 20 PP1c requires the co-crystallisation of PP1c with a diverse array of regulatory subunits and substrates and is beyond the scope of this paper. However, two models to account for this property of regulatory subunits are that these subunits either alter the conformation of PP1c or simply target PP1 to its substrates. Both mechanisms may operate in vivo depending on the regulatory 25 subunits and substrates. For example, evidence for the former model has recently been reported for the enhancement of myosin dephosphorylation by a complex of PP1c and the M_{110} subunit (Johnson et al., 1996, 1997), whereas the enhancement of the dephosphorylation of glycogen phosphorylase and glycogen synthase by the PP1-G_M complex is more consistent with the second

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model (Hubbard and Cohen, 1989).

The identification of the (R/K)(V/I)XF motif also suggests a new approach for determining the physiological roles of PP1-targeting subunits whose functions are unknown. Thus mutation of the (R/K)(I/V)XF motif should disrupt the interaction of many targeting subunits with PP1c without affecting their binding to the target locus. Expression of these mutated proteins under an inducible promoter should lead to displacement of the normal targeting subunit (complexed to PP1c) from its target locus, without disrupting the functions of any other PP1c-targeting subunit complex. Finally, the structural information described here will also facilitate the rational design of drugs that act by disrupting PP1-targeting subunit interactions.

Example 3: Identification of the regions on the M_{110} subunit of protein phosphatase 1M that interact with the M_{21} subunit and with myosin

Abbreviations:- PP1_M, myofibril-associated form of protein phosphatase 1; PP1c, catalytic subunit of protein phosphatase-1; M₁₁₀ and M₂₁, 110 kDa and 21 kDa regulatory subunits of PP1_M; MBP, maltose-binding protein; GST, glutathione-S-transferase.

SUMMARY

We have previously isolated a form of protein phosphatase-1 (PP1_M) from avian smooth muscle myofibrils which is composed of the catalytic subunit of PP1 (PP1c) bound to an M-complex consisting of 110 kDa (M₁₁₀) and 21 kDa (M₂₁) subunits. The interaction of PP1c with an N-terminal region of the M₁₁₀ subunit enhances the dephosphorylation of myosin and suppresses the dephosphorylation of other substrates [Alessi, D.R., MacDougall, L.K., Sola, M.M., Ikebe, M. and Cohen, P. (1992) Eur. J. Biochem 210, 1023-1035;

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Chen, Y.H., Chen, M.X., Alessi, D.R., Campbell, D.G., Shanahan, C., Cohen, P. and Cohen, P.T.W. (1994) FEBS Lett 356, 51-56; Johnson, D.F., Moorhead, G., Caudwell, F.B., Cohen, P., Chen, Y.H., Chen, M.X. and Cohen, P.T.W. (1996) Eur. J. Biochem. 239, 317-325]. In this Example we establish that PP1_M accounts for nearly all the myosin phosphatase activity in myofibrils, that the M_{110} and M_{21} subunits are present at similar concentrations in the myofibrillar fraction and that these subunits are entirely bound to PP1. We demonstrate that the M₂₁ subunit does not interact with PP1c, but with the C-terminal 72 residues of the M₁₁₀ subunit, a region which is 43% identical to residues 87-161 of the M_{21} subunit. A fragment of the M_{21} subunit, M_{21} -(M1-L146), lacking the C-terminal leucine zipper, also bound to the M_{110} subunit, but two other fragments M_{21} -(M1-E110) and M_{21} -(E110-K186) did not. The M_{110} and M_{21} subunits were both found to be myosin-binding proteins. C-terminal 291 residues of the M₁₁₀ subunit, but not the C-terminal 72 residues, bound to myosin, but the N-terminal fragments M_{110} -(M1-E309) and M_{110} -(M1-S477) did not. Thus the region of the M_{110} subunit which stimulates the dephosphorylation of myosin by PP1c is distinct from the region which targets PP1_M to myosin. Remarkably, each myosin dimer was capable of binding about 20 moles of M_{21} subunit and many of the M_{21} -binding sites were located in the myosin "rod domain". The potential significance of this observation is discussed.

Introduction

Protein phosphatase-1 (PP1), one of the major serine/threonine-specific protein phosphatases in eukaryotic cells, is regulated by targetting subunits that direct it to particular subcellular loci, modify its substrate specificity and confer the ability to be regulated by extracellular signals (reviewed in [1, 2]). A significant proportion of the PP1 in vertebrate muscle extracts is associated with myofibrils and has enhanced activity towards the P-light chain of myosin and

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reduced activity towards other substrates, such as glycogen phosphorylase [3, 4]. When isolated from avian (chicken gizzard) [4, 5] or mammalian (pig bladder) [6] smooth muscle, this form of PP1 (PP1_M) was found to be composed of three subunits, namely the catalytic subunit of PP1 (PP1c) and two other proteins with molecular masses of 110 kDa and 21 kDa, termed the M_{110} and M_{21} subunits, respectively [4, 5]. The M_{110} subunit is complexed to both PP1c and the M_{21} subunit [4], and is the component which modulates the substrate specificity of PP1c because selective removal of the M_{21} subunit from PP1_M does not affect the rate at which either myosin or glycogen phosphorylase are dephosphorylated [7].

The M_{110} subunit has been cloned from rat aorta [5], chicken gizzard [8] and rat kidney [9] cDNA libraries. The N-terminus of the M_{110} subunit contains seven ankyrin repeats (residues 39-296 of the rat aorta protein), while alternative splicing in rat uterus [5] gives rise to two different C-termini (Fig 17A), termed Rat1 and Rat2. The C-terminus of Rat1 is virtually identical to the C-terminus of the M_{110} subunit from chicken gizzard (Fig 17A). The sequence of the M_{21} subunit from chicken gizzard is structurally related to the C-terminal region of the M_{110} subunit, the most striking similarities occurring from residues 76-141 of the M_{21} subunit and residues 921-984 of the chicken gizzard M_{110} subunit (54% identity, Fig 17B). However, the C-terminal 53 residues of the M_{21} subunit from chicken gizzard are strikingly similar (83% identity) to the C-terminal 53 residues of the rat aorta sequence, both terminating in a leucine zipper (Fig 17B, [5]).

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Residues 1-309 of the M_{110} subunit from rat aorta, M_{110} -(M1-E309), mimic the intact M_{110} subunit in stimulating the dephosphorylation of myosin and in suppressing the dephosphorylation of glycogen phosphorylase by PP1c, but a slightly shorter construct M_{110} -(D39-E309) (which still contains the seven ankyrin repeats) is unable to modulate the specificity of PP1c [7]. This

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observation led to the finding that the N-terminal 38 residues, M_{110} -(M1-F38), bind to PP1c and enhance the dephosphorylation of myosin. However, M_{110} -(M1-F38) does not suppress the dephosphorylation of glycogen phosphorylase, suggesting that the ankyrin repeats either contain a second PP1c-binding site or prevent glycogen phosphorylase from binding to the active site of PP1c, perhaps by steric hindrance [7].

A 13 residue peptide $G_{M^-}(G63-N75)$ from the subunit (G_M) which targets PP1c to glycogen and the sarcoplasmic reticulum in striated muscle, has been co-crystallised with PP1c and the structure of the complex solved to 3\AA resolution [2]. These studies showed that a hexapeptide sequence in $G_{M^-}(G63-N75)$ (Arg-Arg-Val-Ser-Phe-Ala) (SEQ ID No 3) binds to a small hydrophobic groove on the surface of PP1c forming a β -sheet which runs parallel to another β -sheet in PP1c. Moreover, inspection of other mammalian PP1c-binding proteins reveals that almost all contain an Arg/Lys-Val/Ile-Xaa-Phe motif that is likely to be critical for interaction with PP1c [2]. For example, a Lys-Val-Lys-Phe (SEQ ID No 5) motif is present between residues 35 and 38 of the M_{110} subunit and the deletion of residues 36-38 from M_{110} -(M1-F38) prevents this peptide from stimulating the dephosphorylation of myosin, and from disrupting the interaction of PP1c with other targetting subunits [2].

The finding that a region near the N-terminus of the M_{110} subunit binds to PP1c and modulates its specificity raised the question of which region on the M_{110} subunit interacted with the M_{21} subunit, and how the PP1_M complex is targeted to the myofibrils. In this Example we identify regions near the C-terminus of the M_{110} subunit that interact with the M_{21} subunit and with myosin, and demonstrate that the M_{21} subunit is also a myosin-binding protein. These findings indicate that the domain of the M_{110} subunit which enhances the dephosphorylation of the myosin P-light chain is distinct from the region which targets PP1c to the contractile apparatus.

MATERIALS AND METHODS

Materials

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- PP1_M [4] and the dephosphorylated form of myosin [10] were isolated from chicken gizzard, and the rod-domain and light meromyosin were obtained by subdigestion of chicken gizzard myosin with papain and chymotrypsin, respectively [11]. PP1_G was purified from rabbit skeletal muscle by Dr G. Moorhead in this laboratory [12] and PP1c dissociated from the glycogen-binding subunit by incubation for 2 h in 2 M LiBr and then purified by gel-filtration on a 30 x 1 cm column of Superose 12 (Pharmacia, Milton Keynes, UK) in the presence of LiBr (0.5 M). All other chemicals were from BDH Chemicals (Poole, UK) or Sigma (Poole, UK).
- Construction of vectors for the expression of fragments of the M_{110} subunit from rat aorta (rat2 sequence in Fig 17A) as glutathione-S-transferase (GST) fusion proteins in E. coli.
 - A construct pGEX- M_{110} -(M1-E309) for the expression of GST- M_{110} -(M1-E309) from rat aorta was produced as described previously [7]. A construct for the expression of GST- M_{110} -(M1-S477) was prepared by subcloning a *Xhol-HindIII* fragment (encoding L24-S477) of pKS- M_{110} -(M1-S477) described in [5] into the same sites of pGEX- M_{110} -(M1-E309). The resulting construct expressed a GST- M_{110} -(M1-S477) fusion protein with the additional amino acids SAANSISSLIHRD* (SEQ ID No 27) after S477. An expression construct for GST- M_{110} -(M377-K976) was produced by deleting a *Ncol-Ncol* fragment of the construct pGEX- M_{110} -(L24-K976) [7].
- Construction of vectors for the expression of C-terminal fragments of the M_{110} subunit from chicken gizzard (Ch1 sequence in Fig 17A, [5]) as maltose binding

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protein (MBP) fusion proteins in E. coli.

A pT7.7 vector for the expression of the C-terminal 291 residues of the M₁₁₀ subunit from chicken gizzard, pT7-M₁₁₀-(R714-I1004) was described previously [7]. A construct for the expression of MBP-M₁₁₀-(R714-I1004) was produced by cloning an *NdeI-Bam*HI fragment of pT7-M₁₁₀-(R714-I1004) into the pMAL-HA vector (New England Biolabs). Removal of a *HindIII-HindIII* restriction fragment from pMBP-M₁₁₀-(R714-I1004) allowed expression of MBP-M₁₁₀-(R714-L934) with the sequence GTGRRFTTS (SEQ ID No 28) added to its C-terminus. Removal of a *NdeI-HindIII* restriction fragment from pMBP-M₁₁₀-(R714-I1004), followed by filling in the overhanging ends and religating them, allowed expression of MBP-M₁₁₀-(K933-I1004).

Construction of vectors for the expression in E. coli. of the M_{21} subunit from chicken gizzard [5], with and without the C-terminal leucine zipper domain.

A pT7.7 vector for the expression of the entire coding region (M1-K186) of the M_{21} subunit was described previously [7]. The leucine zipper motif of the M_{21} subunit was deleted by removing a SacI-BamHI restriction fragment from pT7.7 M_{21} , filling in the overhanging ends and religating them. The construct expressed M_{21} -(M1-R144) with an extra I and L after residue 144. The M_{21} -(M1-R144) protein was insoluble when expressed and was purified as described for the expressed M_{21} subunit [7].

Construction of vectors for the expression of the M_{21} subunit from chicken gizzard [5] and fragments of the M_{21} subunit as glutathione-S-transferase (GST) fusion proteins in E. coli.

A construct expressing GST-M₂₁ was produced by inserting a NdeI-HindIII fragment of pT7.7 M₂₁ encoding M1-K186 into the same sites of the pGEX

vector modified to include an *NdeI* site. A construct expressing GST-M₂₁-(M1-E110) plus an additional Ala residue at the C-terminus was constructed by deleting a *XhoI-HindIII* fragment of pGEX-M₂₁, filling in the overhanging ends and religating them. In order to express GST-M₂₁-(E110-K186), a *NdeI-XhoI* restriction fragment of pGEX-M₂₁ was deleted and the overhanging ends filled in and religated.

Expression of proteins in E. coli.

This was carried out essentially as described in [7], except that, after freezing the cells at -80°C and thawing, the lysates were not treated with DNAase but sonicated for 4 min on ice (ensuring that the temperature remained below 4°C) until the suspension was no longer viscous. The soluble GST-fusion proteins and MBP-fusion proteins were purified from the supernatant by affinity chromatography on glutathione-Sepharose (Sigma) and amylose resin (New England Biolabs), respectively, according to the instructions of the manufacturers. After expression in E. coli M₁₁₀-(R714-I1004) was the major soluble protein and all experiments with this fragment were performed using the bacterial extracts.

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The chicken gizzard M₂₁ subunit was isolated from *E. coli* extracts as described [7]. M₂₁ subunit lacking the leucine zipper domain, M₂₁-(M1-L146), like the M₂₁ subunit itself, was obtained in inclusion bodies and therefore recovered in the pellet obtained after centrifugation of the bacterial lysates for 30 min at 28 000 x g. The inclusion bodies were washed three times in 50 mM Tris/HCl pH 7.5, 0.1M NaCl, 10 mM EDTA, 0.1% (by vol) 2-mercaptoethanol, 1 mM benzamidine, 0.2 mM phenylmethylsulphonyl fluoride and 0.5% (by mass) Triton X-100, then resuspended in 50 mM Tris/HCl pH 7.5, 1 mM EDTA, 1 mM EGTA, 0.03% (by mass) Brij-35, 0.1% (by vol) 2-mercaptoethanol. An aliquot (containing 3 mg protein) was made 0.5% (by vol) in trifluoroacetic

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acid, sonicated, centrifuged for 2 min at 13,000 x g and the supernatant (containing the solubilised M₂₁ subunit) loaded on to a Vydac C18 column (Separations Group, Hesperia, CA, USA) equilibrated in 0.1% (by vol) trifluoroacetic acid. The column was developed with a linear acetonitrile gradient at a flow rate of 1.0 ml / min with an increase in acetonitrile concentration of 1% per min. Homogeneous M₂₁ subunit, which eluted at 42% acetonitrile, and M₂₁-(M1-L146) which eluted at 40% acetonitrile were dried in a vacuum concentrator redissolved in water, redried and then dissolved in 50 mM Tris/HCl pH 7.5, 0.1 mM EGTA, 0.03% (by mass) Brij-35, 0.1% (by vol) 2-mercaptoethanol.

Removal of GST and MBP tags from fusion proteins.

GST-M₁₁₀-(1-477) was cleaved with thrombin and the proteinase removed using benzamidine agarose [7]. GST- M_{21} -(E110-K186) (1mg/ml) was cleaved 15 by incubation for 1 h at 30°C with 10 μ g/ml thrombin, while GST-M₂₁-(M1-E110) (1mg / ml) was cleaved by incubation for 3 h at 30°C with 1 μ g/ml thrombin, because it was more susceptible to degradation by thrombin. MBP- $M_{110}(K933-I1004)$ (1 mg / ml) was cleaved by incubation for 8 h at 23°C with Factor Xa (10 μ g/ml). Other conditions and removal of Factor Xa were carried out as described for thrombin [7].

Preparation of phosphorylated myosin P-light chain and phosphatase assays.

³²P-labelled myosin P-light chains containing 1.0 mol phosphate per mol 25 subunit was prepared by phosphorylation with smooth muscle myosin light chain kinase [4]. The dephosphorylation of myosin P-light chain (1 μ M) was carried out as in [4] and one unit of activity (U) was that amount which catalysed the release of 1 μ mole of phosphate in one min. When assaying PP1_M in immunoprecipitates from the myofibrillar extracts, the tubes were 30

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shaken continuously and 3 nM okadaic acid was included to inhibit any PP2A present.

Immunoprecipitation of PPI_M from myofibrillar extracts.

Antibodies raised against the PP1_M holoenzyme (1 μ g), which recognise both the M_{110} and M_{21} subunits, but not PP1c, affinity purified antibodies specific for either the M_{110} subunit or M_{21} subunit (5 μ g) [7], and control IgG (5 μ g) were conjugated separately to 10 µl of pelleted protein G-Sepharose. After incubation for 30 min at 4°C, the Protein G-Sepharose-antibody conjugate was washed three times with 50 mM Tris/HCl pH 7.5, 0.1 mM EGTA, 0.03% (by mass) Brij-35, 0.3M NaCl, 0.1% (by vol) 2-mercaptoethanol before addition of a 100 μ l of myofibrillar extract (prepared as in [4]) which had been diluted 10-fold in 50 mM Tris/HCl pH 7.5, 0.1 mM EGTA, 0.1% (by vol) 2mercaptoethanol, 0.2 mM phenylmethylsulphonyl fluoride, 1 mM benzamidine, 10 μg/ml leupeptin containing 1 mg/ml bovine serum albumin. incubation for 1 h at 4°C, with shaking, a 10 µl aliquot of the suspension was removed to measure the total activity. The remaining 90 µl was centrifuged for 1 min at 13,000 x g, the supernatant was removed, and the pellet washed twice in dilution buffer containing 0.2 M NaCl and 0.03% (by mass) Brij-35 (but no bovine serum albumin), once in dilution buffer and then resuspended in 90 μ l of dilution buffer. Myosin P-light chain phosphatase activity was then measured in the supernatant and the resuspended pellet at a further 30-fold final dilution.

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Myosin binding assays. Myosin $(0.5 \text{ mg} / \text{ml}, 1 \,\mu\text{M} \text{ in terms of myosin heavy})$ chains) in 10 mM Hepes pH 7.5, 50 mM KCl, 5 mM MgCl2, 0.1% (by vol) 2-mercaptoethanol, was mixed with PP1_M, M₂₁ subunit, or fragments of the M₁₁₀ and M₂₁ subunits, at the concentrations indicated in figure legends. After incubation for 15 min at 0°C, the solutions were centrifuged for 2 min at

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13,000 x g, the supernatants removed, and the pellets washed twice in 10 mM Hepes pH 7.5, 50 mM KCl, 5 mM MgCl2, 0.1% (by vol) 2-mercaptoethanol before resuspension in 50 mM Tris-HCl pH 7.5, 0.1 mM EGTA, 0.03% (by mass) Brij 35, 0.6 M NaCl, 0.1% (by vol) 2-mercaptoethanol. Aliquots of the supernatant, the resuspended pellet and the suspension before centrifugation were either assayed for myosin P-light chain phosphatase activity or denatured in SDS and analysed by SDS/polyacrylamide gel electrophoresis.

Preparation of a complex between GST- M_{21} and M_{110} -(R714-I1004).

GST- M_{21} (10 μ g) was mixed with 80 μ l of bacterial extract expressing M_{110} -(R714-I1004). After incubation for 15 min at ambient temperature the solution was added to 20 μ l (packed volume) of glutathione-Sepharose equilibrated in 50 mM Tris HCl pH 7.5, 0.1 mM EGTA, 0.03% (by mass) Brij 35, 0.1% (by vol) 2-mercaptoethanol, 0.2 mM phenylmethylsulphonyl fluoride, 1 mM benzamidine and 0.15 M NaCl. After incubation for 30 min at 4°C with shaking, the Sepharose was washed three times in the same buffer before eluting the complex with buffer containing 20 mM glutathione pH 8.0.

20 Other procedures.

Proteins were labelled with digoxigenin and Far Western analyses carried out as described [4], except that the digoxigenin-labelled probe was used at a concentration of 0.2 μ g/ml instead of 2 μ g/ml. SDS/polyacrylamide gel electrophoresis was carried out on 7.5-15% gels according to Laemmli [13] and on 16.5% gels according to Schagger and von Jagow [14]. Protein was estimated according to Bradford [15].

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Results

 $PP1_{M}$ accounts for nearly all the myosin phosphatase activity in extracts prepared from chicken gizzard myofibrils.

80-90% of the myosin phosphatase activity present in chicken gizzard homogenates is recovered in the myofibrils [4]. In the present study, we used antibodies that recognise the M_{110} and/or the M_{21} subunits of chicken gizzard PP1_M [7] to immunoprecipitate the myosin P-light chain phosphatase activity from the myofibrillar extracts. About 90% of the activity was immunoprecipitated by antibodies raised against the PP1_M holoenzyme (Fig 18A) which recognise both the M_{110} and M_{21} subunits in immunoblotting experiments, but not PP1c. Similarly, about 80% of the myosin P-light chain phosphatase activity in the myofibrillar extracts was immunoprecipitated by either the anti- M_{110} antibody or by the anti- M_{21} antibody (Fig 18A). Thus, most of the myosin P-light chain phosphatase activity in myofibrillar extracts is catalysed by PP1c present as a complex containing both the M_{110} and the M_{21} subunits.

Immunoblotting experiments demonstrated that the ratio M_{110} : M_{21} in myofibrillar extracts was identical to the ratio of these subunits in purified PP1_M (Fig 18B), which is 1:1 [4]. These immunoblotting experiments also demonstrated that PP1_M comprises 0.1% of the protein in the myofibrillar extract (see legend to Fig 18B), identical to the proportion estimated from the fold-purification needed to obtain pure PP1_M from this fraction (see Table 1 in Ref 4). These experiments imply that PP1_M accounts for virtually all the myosin phosphatase activity associated with myofibrils, and that neither the M_{110} nor the M_{21} subunit is present in a significant molar excess over PP1c in the myofibrils.

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Identification of a region on the M_{110} subunit that binds to the M_{21} subunit.

PP1_M and several fragments of the M₁₁₀ subunit, were subjected to SDS/polyacrylamide gel electrophoresis (Fig 19A) and, after transfer to nitrocellulose, the blots were probed with digoxigenin-labelled M₂₁ subunit (Fig 19B). These experiments showed that the M₂₁ subunit recognised the full length M₁₁₀ subunit (Fig 19B, track 1), M₁₁₀-(R714-I1004) (Fig 19B, tracks 2 and 3) and M₁₁₀-(K933-I1004) (Fig 19B, track 5), but not M₁₁₀-(R714-L934) (Fig 19B, track 4), M₁₁₀-(M1-E309) (Fig 19B, track 7) or M₁₁₀-(M1-S477) (Fig 19B, track 8). Thus, the M₂₁ subunit binds to the C-terminal 72 residues of the M₁₁₀ subunit. The specificity of this interaction was indicated by the observation that digoxigenin-labelled M₂₁ subunit recognised only M₁₁₀-(R714-I1004) and no other protein in the *E. coli* extract (track 2 in Figs 19A and 19B), nor did it recognise the MBP or GST tags, PP1c (Figs 19A and 19B) or any of the molecular mass markers (data not shown).

Consistent with the results in Fig 19, digoxigenin-labelled MBP- M_{110} -(R714-I1004) (data not shown) and MBP- M_{110} -(K933-I1004) (Fig 20B), but not digoxigenin-labelled MBP- M_{110} -(R714-L934) (data not shown), recognised the full length M_{21} subunit and M_{21} (M1-L146) in Far Western blotting experiments.

The region of the M_{21} subunit that interacts with the M_{110} subunit.

Digoxigenin-labelled M₂₁-(M1-L146) recognised the same fragments of the M₁₁₀ subunit as the full length M₂₁ protein (Fig 19C), demonstrating that the C-terminal leucine zipper of the M₂₁ subunit is not required for interaction with the M₁₁₀ subunit. However, neither digoxigenin-labelled GST-M₂₁-(M1-E110) nor digoxigenin-labelled GST-M₂₁-(E110-K186) recognised M₁₁₀-(K933-I1004) in Far Western blotting experiments (data not shown). Consistent with these findings, digoxigenin-labelled M₁₁₀-(K933-I1004) recognised the full length M₂₁

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protein and M_{21} -(M1-L146), but not M_{21} -(M1-E110) or M_{21} -(E110-K186) in Far Western blotting experiments (Figs 20A and B). However, digoxigenin-labelled M_{110} -(K933-I1004) did recognise a proteolytic fragment of the M_{21} subunit with an apparent molecular mass only slightly larger than M_{21} -(M1-E110) (Fig 20B, track 2 and compare tracks 2 and 4 in Fig 20A). These results are considered further under Discussion.

The isolated M_{21} subunit dimerizes and the region involved in dimerization is identical to that which interacts with the M_{110} subunit.

Although the M_{110} subunit binds to both PP1c and the M_{21} subunit [4], and removal of the M_{21} subunit does not alter the specificity of the PP1_M complex [7], an interaction between the M_{21} subunit and PP1c had not been excluded. In order to examine this point, PP1c and the M_{21} subunit were mixed together and subjected to gel filtration on Superose 12. The M_{21} subunit eluted just before the 37 kDa PP1c protein, demonstrating that they do not form a high affinity complex and suggesting that the isolated M_{21} subunit dimerizes (data not shown). These results were supported by the finding that digoxigenin-labelled full length M_{21} subunit recognised the M_{21} subunit as well as the M_{110} subunit, but not PP1c, in Far Western blotting experiments (Fig 21, track 1). Similar results were obtained with M_{21} -(M1-L146) (Fig 21, track 2). Digoxigenin-labelled M_{21} subunit, like digoxigenin-labelled M_{110} -(K933-I1004), recognised a fragment of the M_{21} subunit that migrated slightly more slowly than M_{21} -(M1-E110), but did not recognise M_{21} -(M1-E110) or M_{21} -(E110-K186) (Tracks 2, 4 and 5 in Figs 20B and 20C).

Identification of a region on the M_{110} subunit that binds to myosin.

When PP1_M (30 nM) was mixed with chicken gizzard myosin (1 μ M) and centrifuged to pellet the myosin, 85% of the myosin P-light chain phosphatase

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was recovered in the pellet (Figs 22 and 23A). In contrast, neither PP1c (Fig 22) nor bovine serum albumin (data not shown) bound to myosin under these conditions. After removal of the M_{21} subunit from PP1_M [7], the M_{110} -PP1c complex (PP1_M(ΔM_{21}) still pelleted with myosin in a similar manner to PP1_M itself (Fig 22), indicating that the M_{110} subunit is a myosin-binding protein.

In order to identify the myosin-binding domain(s), several fragments of the M_{110} subunit were expressed and purified from $E.\ coli$ extracts and their binding to myosin was studied. GST- M_{110} -(M1-S477), like GST- M_{110} -(M1-E309) [7], stimulated the PP1c-catalysed dephosphorylation of the myosin P-light chain and inhibited the dephosphorylation of glycogen phosphorylase in a similar manner to the full length M_{110} subunit (data not shown). However, neither GST- M_{110} -(M1-S477) nor GST- M_{110} -(M1-E309) bound to myosin (data not shown), even after removal of the GST-tag from GST- M_{110} -(M1-S477) (Fig 23A).

A fragment comprising GST- M_{110} -(M377-K976) from rat aorta migrated as multiple bands on SDS/polyacrylamide gels after purification on glutathione-Sepharose (Fig 23A), indicating cleavage at multiple sites within the M_{110} subunit. Only the largest fragment, with an apparent molecular mass corresponding to undegraded GST- M_{110} -(M377-K976) bound to myosin (Fig 23A), suggesting that the myosin binding site(s) was located towards the C-terminus of the M_{110} subunit. Consistent with this finding, M_{110} -(R714-I1004) from chicken gizzard also bound to myosin (Fig 23B). However, M_{110} -(K933-I1004), which bound to the M_{21} subunit (Fig 20B), did not bind to myosin in these experiments (Fig 23B).

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The M_{21} subunit, and a complex between M_{21} and M_{110} (R714-I1004) bind to myosin.

After purification on glutathione-Sepharose, GST-M₂₁ migrated as four protein staining bands (track 1 in Fig 20A), the two species of highest apparent molecular mass being recognised by the anti-M₂₁ antibody (Fig 23B). The apparent molecular mass of the slowest migrating band (47 kDa) corresponds to undegraded GST-M₂₁ and this species bound to myosin (Fig 23B). The next most slowly migrating band had an apparent molecular mass of 38 kDa, slightly less than that of GST-M₂₁-(M1-E110) (data not shown) indicating that it corresponds to GST fused to about the first 100 residues of the M₂₁ subunit; this fragment hardly bound to myosin (Fig 23B).

Bacterial extracts expressing M_{110} -(R714-I1004) were mixed with GST- M_{21} and the resulting complex was purified on glutathione-Sepharose. This complex bound quantitatively to myosin (Fig 23B). In contrast, the GST- M_{21} fragment of apparent molecular mass 38 kDa was not complexed to M_{110} -(R714-I1004) and did not bind to myosin (Fig 23B). The C-terminal fragment of the M_{21} subunit, M_{21} -(E110-K186) also did not bind to myosin under these conditions (data not shown).

Multiple binding sites for the M_{21} subunit on the myosin molecule.

The molar ratio myosin: PP1_M in chicken gizzard is about 80:1 in vivo [4] and the myosin binding experiments described above were therefore carried out using a large (ten fold) molar excess of myosin over either the M_{21} or the M_{110} subunit. However, further experiments carried out with the M_{21} subunit in excess revealed that, remarkably, 20 or more moles of M_{21} subunit could be bound to each myosin dimer (Fig 24A). Many of the binding sites were located in the region of myosin involved in filament formation, because the M_{21}

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subunit was pelleted with the myosin "rod" domain even when the molar ratio M_{21} : myosin dimer was 10:1 (Fig 24B). A shorter portion of the rod, termed light meromyosin, also bound the M_{21} subunit avidly. However, a fragment of the M_{21} subunit lacking the first 15 residues from the N-terminus, which was a contaminant in this preparation, did not bind to light meromyosin (Fig 24B), although it bound to the longer myosin rod (Fig 24B). The M_{21} subunit lacking the C-terminal leucine zipper, M_{21} -(M1-L146), bound to both myosin and the rod domain, but fewer moles of M_{21} -(M1-L146) could be bound and this C-terminally truncated species did not bind to light meromyosin under the conditions studied (Fig 24C).

Multiple forms of the M_{110} subunit

Comparison of two different clones encoding the M_{110} subunit from chicken gizzard revealed a 123 bp (41 amino acid) deletion/insertion after Asn-511 (Fig. 17, [8]). Since the rat aorta sequence [5] showed considerable variation from the chicken sequences in this region, compared to the high degree of sequence similarity throughout most of the rest of the molecule (Fig 17), it seemed probable that forms of the rat M₁₁₀ subunit also existed that varied in this middle section of the protein. PCR of the "variable region" of several rat aorta clones gave fragments of either 608 bp or 776 bp. Direct sequencing of these fragments showed an in frame insertion of 168bp (56 amino acids) after Ser-552 (Fig. 1); i.e. a slightly different position from the deletion reported for the chicken gizzard M₁₁₀ subunit (Fig 17). Furthermore, a different 62 amino acid deletion/insertion in this section is apparent by comparison of the rat aorta sequences with that of the M_{110} protein from rat kidney (Fig. 1) [9]. While it is likely that most of these variations arise by alternative splicing of the mRNA, Southern blotting of rat genomic DNA revealed the presence of two closely related genes (data not shown).

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Discussion

The contraction of smooth muscle is triggered by phosphorylation of the P-light chain of myosin catalysed by myosin light chain kinase. However, the identity of the myosin P-light chain phosphatase remained unclear for many years. In 1992 we reported that 80-90% of the myosin phosphatase activity in chicken gizzard homogenates was associated with myofibrils and purified a myosin phosphatase to homogeneity from this fraction [4]. This enzyme, termed PP1_M, was found to be composed of the β -isoform of PP1c (termed the δ -isoform in [16]) and an "M-complex" consisting of two other subunits [4] whose molecular masses were 21 kDa (M_{21}) [5] and 110 kDa (M_{110}) [5, 8], respectively. Further evidence that a form of PP1 was the major myosin phosphatase in smooth muscle was indicated by the finding that tautomycin (a much more potent inhibitor of PP1 than PP2A [17]) stimulated the contraction of permeabilised mammalian smooth muscle fibres at much lower concentrations than okadaic acid [18] (a much more potent inhibitor of PP2A than PP1 [19]).

Two further pieces of evidence presented in this Example establish that $PP1_M$ accounts for most, if not all, of the myosin phosphatase activity associated with chicken gizzard myofibrils, reinforcing the view that it is likely to be the major myosin P-light chain phosphatase in vivo. Firstly, nearly all the myosin P-light chain phosphatase activity was immunoprecipitated by antibodies that recognise either the M_{110} or the M_{21} subunit specifically (Fig 18A). Secondly, $PP1_M$ was found to represent 0.1% of the protein in the myofibrillar extracts whether its concentration was calculated from the increase in specific activity needed for purification to homogeneity [4] or from immunoblotting experiments with the anti- M_{110} and anti- M_{21} antibodies (Fig 18B). Had another enzyme been the major myosin phosphatase in the myofibrillar extracts the enrichment estimated by immunoblotting with anti- M_{110} and anti- M_{21} antibodies would have been

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much higher.

The experiments presented in Fig 18 also demonstrate that the M_{110} and M_{21} subunits are not present in myofibrillar extracts in a significant molar excess over PP1c and that all the M_{110} subunits are complexed to M_{21} subunit and vice versa. The M₂₁ subunit was found to bind to the C-terminal 72 residues of the chicken gizzard M_{110} subunit (Figs 19 and 25), a region whose amino acid sequence is 43% identical to residues 87-161 of the M₂₁ subunit (Fig 17B). The C-terminal leucine zipper of the M₂₁ subunit (residues 145-186) is not required for interaction with the M₁₁₀ subunit, and the site on the M₂₁ subunit which interacts with the M₁₁₀ subunit lies within about the N-terminal 120 residues (Fig 20B). Interestingly, the same region is essential for the dimerisation of the M₂₁ subunit (compare Figs 20B and 20C), suggesting that the region(s) involved in interaction is probably located between residues 60 and 120 of the M_{21} subunit and 906-965 of the M_{110} subunit from chicken gizzard; i.e. the regions with greatest amino acid identity between these two proteins (Fig 17). More digoxigenin-labelled M21 subunit bound to the M110 subunit than to the M_{21} subunit in Far Western blotting experiments (Fig 21), consistent with the observation that M_{110}/M_{21} heterodimers form in vivo, but not M_{21}/M_{21} homodimers. The finding that the C-terminus of the M_{110} subunit interacts with the M₂₁ subunit explains why preparations of PP1_M comprising PP1c complexed to N-terminal fragments of the M₁₁₀ subunit do not contain the M_{21} subunit [8, 20].

PP1_M binds to the dephosphorylated form of myosin and our data demonstrate that the M₁₁₀ subunit (Fig 22) and the M₂₁ subunit (Fig 23B and Fig 24) are both myosin-binding proteins. The C-terminal 600 residues of the M₁₁₀ subunit from rat aorta, M₁₁₀-(M377-K976) (Fig 23A) and the C-terminal 291 residues of the M₁₁₀ subunit from chicken gizzard, M₁₁₀-(R714-I1004), bound to myosin, but the C-terminal 72 residues of the M₁₁₀ subunit, M₁₁₀-(K933-1004), did not

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(Fig 23B), indicating that a myosin-binding domain is likely to be situated in the M_{110} subunit just N-terminal to the M_{21} -binding domain (Fig 25). In contrast, two N-terminal fragments of the M_{110} subunit M_{110} -(M1-S477) (Fig 23A) and M_{110} -(M1-E309) (data not shown) did not bind to myosin under the conditions studied. Since M_{110} -(M1-E309) [7] and M_{110} -(M1-S477) (data not shown) stimulate the dephosphorylation of myosin and inhibit the dephosphorylation of glycogen phosphorylase by PP1c, and in a similar manner to full length M_{110} subunit, these results show that the region of the M_{110} subunit which stimulates the dephosphorylation of the myosin P-light chain is distinct from that which binds the dephosphorylated form of myosin and thereby targets PP1_M to the contractile apparatus.

Digestion of chicken gizzard PP1_M with chymotrypsin cleaves the M₁₁₀ subunit to a fragment with an apparent molecular mass of 58 kDa and a form of PP1, termed here PP1_M*, can then be isolated by gel-filtration which appears to comprise just the 58 kDa fragment and PP1c in a 1:1 molar ratio [8]. The 58 kDa fragment, like the M₁₁₀ subunit, has a blocked N-terminus and seven tryptic peptides isolated were located between residues 286 and 467, suggesting that the 58 kDa fragment corresponds to the N-terminal portion of the M₁₁₀ subunit [8]. PP1_M* was reported to bind to myosin, albeit less effectively than PP1_M [8], suggesting the presence of a myosin-binding domain within the 58 kDa fragment. This result is in apparent conflict with the present study, also migrates which fragment M_{110} -(M1-S477), the SDS/polyacrylamide gels with an apparent molecular mass of 58 kDa, did not bind to dephosphorylated myosin under conditions where 80-90% of the $PP1_M$ and M_{110} -(R714-I1004) was pelleted with myosin (Fig 23A). One possible explanation for this discrepancy is that PP1_M* also contains small myosinbinding fragments from the C-terminus of the M₁₁₀ subunit which still interact with the N-terminal 58 kDa fragment, but are too small to be detected by SDS/polyacrylamide gel electrophoresis. In a separate study heavy meromyosin (50 μ g) was found to bind partially to 2 mg of M₁₁₀-(1-633) coupled to Affigel 15, at very low ionic strength but not at 150-200 mM NaCl [21]. The significance of this observation is unclear because of the extremely high concentration of the M₁₁₀-(1-633) used in these experiments. The average intracellular concentration of PP1_M in chicken gizzard is about 1 μ M, 100-fold lower than the concentration of myosin. In the present study, we analysed the binding of the M₁₁₀ subunit and its subfragments (30-100 nM) to myosin (1 μ M) using low concentrations of these proteins to try and ensure that only high affinity binding sites were identified.

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The isolated M_{21} subunit also bound to myosin and up to 20 moles of M_{21} subunit could be bound to each myosin dimer (Fig 24). These observations indicate that each myosin molecule contains multiple binding sites for the M_{21} subunit, many of which are located within the "rod domain" (Figs 24B and 24C). In vivo, the molar ratio $PP1_{M}$: myosin is about 1:80 and yet, during muscle relaxation, all the myosin P-light chains can be dephosphorylated by $PP1_{M}$ within seconds. This implies that $PP1_{M}$ must be highly mobile within the myofibrils and move extremely rapidly from one myosin molecule to another. The "off rates" for binding of $PP1_{M}$ to myosin must therefore be very fast as well as the "on rates". It is tempting to speculate that the presence of multiple binding sites on myosin for the M_{21} subunit (and perhaps for the M_{110} subunit as well) allows $PP1_{M}$ to "slide" rapidly from one myosin molecule to another.

Example 4: Design of small molecules to modulate the properties of PP1

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Table A is a print-out of the atomic coordinates of the protein phosphatase-1 peptide coordinates as deduced in Example 2. The format is Protein Data Bank. The structure of the protein phosphatase-1 catalytic subunit (PP1c) in complex with a 13-residue peptide (G_M peptide) corresponding to a site of interaction between PP1c and the glycogen targeting subunit provides a frame-

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work for the rational design of small molecules to modulate the functions and properties of PP1 in vivo. Knowledge of the structural nature of the interactions between the G_M peptide and PP1c allows the design of inhibitors that mimic these interactions. These inhibitors may be designed for increased potency, cell permeability and with improved pharmokinetic properties.

Computer graphics systems may be used to design such inhibitors in conjunction with computer graphics modelling software such as SYBIL available from: Tripos Inc, 16995 S Hanley Road, St Louis, Missouri 63144-2913, USA and LUDI available from: Molecular Simulations Inc, 9685 Scranton Road, San Diego, CA 92121-3752, USA, and in conjunction with the atomic coordinates shown in Table A.

Example 5: Effect of peptide derived from p53BP2 binding site to PP1 in vivo

The function of p53BP2 is ascertained by examining the *in vivo* effect of peptides based on the sequence of the p53BP2 binding site to PP1. This may be done by reference to the consensus peptide sequence described in the previous Examples and by reference to the crystal structure in Example 2. The peptide is introduced into cultured cells using penetratin available from Appligene. Other importins may also be used. Alternatively cDNA specifying p53BP2 proteins mutant in the p53BP2 binding site to PP1 are transfected in cultured cells. The effect of these agents on the cell cycle and apoptosis are assessed by a number of methods, for example WAF1 ELISA and Nuclear Matrix Protein ELISA assays (Amersham).

The effect of the p53BP2 peptide is to modulate the interaction between PP1 and p53BP2 in vivo and affect cell regulation and apoptosis. The p53BP2 peptide may also be micro-injected into the cell.

1 N LYS 3 CA LYS 4 CB LYS 5 CG LYS 6 CD LYS 7 CE LYS 8 NZ LYS 12 C LYS 13 O LYS 14 N LEU 16 CA LEU 17 CB LEU 18 CG LEU 19 CD1 LEU 20 CD2 LEU 21 C LEU 22 O LEU 23 N ASN	66666666677777778	Table A -10.263 -9.182 -9.220 -10.284 -9.809 -8.832 -7.498 -7.814 -6.854 -7.746 -6.527 -6.840 -5.670 -4.775 -6.186 -5.892 -6.497 -4.656	46.372 46.177 47.277 47.095 46.200 46.919 47.216 46.1624 46.861 47.599 47.782 48.881 48.121 45.594 47.782 48.424	91.126 90.159 89.092 88.015 86.868 85.927 86.540 90.8303 92.005 92.800 94.141 95.106 94.589 96.496 93.063 93.6675 92.627	1.00 53.07 1.00 53.07 1.00 45.20 1.00 45.20 1.00 45.20 1.00 45.20 1.00 45.20 1.00 45.20 1.00 42.33 1.00 43.14 1.00 24.45 1.00 18.34 1.00 22.21 1.00 42.81 1.00 23.55 1.00 17.03	000000000000000000000000000000000000000
25 CA ASN 26 CB ASN 27 CG ASN 28 OD1 ASN 29 ND2 ASN 32 C ASN 33 O ASN 34 N ILE 36 CA ILE 37 CB ILE 38 CG2 ILE 39 CG1 ILE 41 C ILE 41 C ILE 41 C ILE 42 O ILE 41 C ILE 42 O ASP 46 CB ASP 46 CB ASP 47 CG ASP 48 OD1 ASP 49 OD2 ASP 50 C ASP 51 O ASP 51 O ASP 52 N SER 55 CB SER 56 OG SER 58 C SER 59 O SER 56 OG SER 58 C SER 56 OG SER 56 CB ILE 66 CD1 ILE 67 C ILE 66 CD1 ILE 67 C ILE 66 CD1 ILE	8 8 8 8 8 9 9 9 9 9 9 9 9 9 9 10 0 10 0	-4.000 -3.204 -3.4843 -3.46439 -3.110 -3.9016 -3.9155 -3.1742 -3.865 -0.756 -0.8334 -2.126 0.426 0.2487 -0.925 1.5345 -0.964 3.1689 1.5345 -0.8331 -0.8499 -1.501 3.6887 -0.764	44.156 43.744 42.312 41.503 44.5100 43.828 44.5100 43.828 44.837 42.839 44.837 42.839 41.886 41.8	91.610 91.1063 91.0683 91.0993 91.0993 91.0993 92.0280 93.0282 93.0	1.00 18.54 1.00 15.30 1.00 14.53 1.00 7.61 1.00 12.23 1.00 12.56 1.00 14.41 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 45.77 1.00 46.34 1.00 50.94 1.00 55.84 1.00 2.00 1.00 43.44 1.00 2.00	000000000000000000000000000000000000000
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ATOM ATOM	175	Cλ	LYS	23	20.828	47.458	99.740	1.00 49.50	0
ATOM	176	CB	LYS	23	21.565	48.471	98.852	1.00 94.25	0
ATOM	177	CG	LYS	23	20.639	49.422	98.085	1.00 59.71 1.00 94.25	0
MOTA	178	CD	LYS	23	21.341	50.716 51.775	97.688 97.214	1.00 59.77	Ö
MOTA	179	CE	LYS	23	20.346 19.448	52.240	98.306	1.00 59.94	Ö
MOTA	180	NZ	LYS LYS	23 23	19.739	48.190	100.528	1.00 49.47	Ŏ
MOTA	184 185	С О	LYS	23	18.659	48.488	99.998	1.00 59.94	ō
MOTA MOTA	186	N	PRO	24	19.990	48.458	101.B13	1.00 37.72	0
ATOM	187	CD	PRO	24	21.185	48.148	102.614	1.00 2.00	0
ATOM	188	CA	PRO	24	18.987	49.165	102.613	1.00 36.73	0
MOTA	189	CB	PRO	24	19.618	49.217 49.208	104.004	1.00 2.00 1.00 2.00	0
MOTA	190	CC	PRO	24 24	21.109 18.798		102.019	1.00 33.26	ŏ
ATOM	191 192	CO	PRO PRO	24	19.752	51.325	101.938	1.00 2.00	ŏ
ATOM ATOM	193	N	GLY	25	17.579	50.835	101.575	1.00 22.79	0
ATOM	195	CA	GLY	25	17.275	52.119	100.981	1.00 22.84	0
ATOM	196	C	GLY	25	16.653	51.904	99.624	1.00 18.00	0
MOTA	197	0	GLY	25	16.098	52.827 50.679	99.037 99.116	1.00 28.24 1.00 2.00	0
MOTA	198	N	LYS	26 26	16.750 16.174	50.351	97.817	1.00 2.00	ŏ
ATOM	200 201	CA CB	LYS LYS	26	16.469	48.892	97.458	1.00 45.45	ō
ATOM ATOM	201	œ	LYS	26	15.931	48.437	96.110	1.00 55.13	0
ATOM	203	CD	LYS	26	16.209	49.435	94.979	1.00 59.31	0
ATOM	204	CE	LYS	26	17.694	49.644	94.691	1.00 60.93	0
MOTA	205	NZ	LYS	26	17.883	50.619 50.624	93.569 97.856	1.00 68.30 1.00 2.00	0
MOTA	209	C	LYS	26 26	14.674 13.916	49.964	98.566	1.00 42.74	ŏ
MOTA	210 211	O N	LYS ASN	27	14.278	51.648	97.111	1.00 2.00	Ö
MOTA MOTA	213	CA	ASN	27	12.894	52.086	97.027	1.00 2.00	0
MOTA	214	CB	ASN	27	12.836	53.526	96.517	1.00 50.37	0
MOTA	215	CG	ASN	27	13.257	54.525	97.563 98.7 4 Q	1.00 56.29 1.00 61.45	0
MOTA	216		ASN	27	12.929 13.982	54.381 55.551	97.142	1.00 59.50	ŏ
ATOM	217	ND2 C	asn asn	27 27	11.964	51.219	96.183	1.00 2.00	Ō
ATOM ATOM	220 221	0	ASN	27	12.384	50.256	95.540	1.00 54.11	0
ATOM	222	N	VAL	28	10.689	51.590	96.209	1.00 12.34	0
MOTA	224	CA	VAL	28	9.646	50.910	95.473	1.00 12.71 1.00 2.00	0
MOTA	225	CB	VAL	28	9.126. 8.777	49.693 50.111	96.283 97.684	1.00 2.00	ŏ
MOTA	226 227	CG1 CG2		28 28	7.932	49.053	95.599	1.00 2.00	0
MOTA MOTA	228	C	VAL	28	8.549	51.935	95.145	1.00 19.22	0
ATOM	229	ŏ	VAL	28	7.757	52.363	96.000	1.00 2.00	0
ATOM	230	N	GLN	29	8.548	52.372	93.892	1.00 26.36 1.00 27.97	0
MOTA	232	CA	GLN	29	7.586	53.365 54.239	93.424 92.325	1.00 27.97 1.00 11.00	Ö
MOTA	233	CB	GLN	29 29	8. 203 7. 47 9	55.543	92.080	1.00 8.47	Ō
MOTA MOTA	234 235	CG CD	GLN	29	7.684	56.541	93.201	1.00 11.72	0
ATOM	236		GLN	29	7.097	57.619	93.198	1.00 13.39	0
ATOM	237	NE2	GLN	29	8.525	56.195	94.159	1.00 12.03 1.00 27.73	0
MOTA	240	C	GLN	29	6.347	52.688 51.934	92.887 91.926	1.00 27.73	ŏ
MOTA	241	0	GLN	29 30	6. 401 5. 22 9	52.941	93.531	1.00 2.00	Ō
MOTA MOTA	242 244	N CA	LEU	30	3.978	52.359	93.087	1.00 2.00	0
MOTA	245	CB	LEU	30	3.157	51.859	94.279	1.00 18.06	0
MOTA	246	CG	LEU	30	3.381	50.416	94.729	1.00 18.06	0
MOTA	247	CD1	LEU	30	4.857	50.152	94.928	1.00 18.06	0
MOTA	248	CD2		30	2.603	50.171 53.441	96.009 92.348	1.00 18.06 1.00 2.00	Ö
MOTA	249	C	LEU	30 30	3.223 3.363	54.621	92.664	1.00 18.06	Ö
MOTA MOTA	250 251	и	LEU GLN	31	2.441	53.050	91.355	1.00 75.78	0
ATOM	253	CA	GLN	31	1.679	54.026	90.599	1.00 80.15	0
ATOM	254	CB	GLN	31	0.782	53.336	89.593	1.00 2.00	0
MOTA	255	ĊG	GLN	31	1.448	52.204	88.883	1.00 2.00 1.00 2.00	0
MOTA	256	CD	GLN	31	0.498	51.469 50. 69 4	87.976 87.122	1.00 2.00	o
MOTA	257		GLN	31	0.933	51.698	88.150	1.00 2.00	Ö
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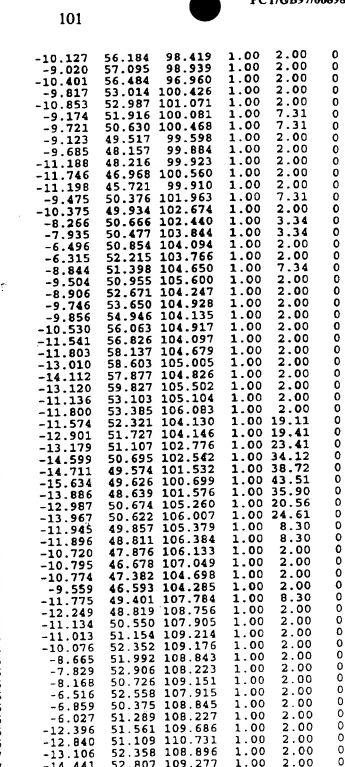
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MOTA	311	CZ	ARG	36 36	-7.726 -8.729	59.383 58.618	94.491 94.064	1.00 22.93 1.00 22.80	0
MOTA MOTA	312 315		ARG ARG	36	-7.987	60.518	95.134	1.00 19.87	0
MOTA	318	C	ARG ARG	36 36	-5.394 -6.415	55.005 55.305	96.6 5 7 97. 27 9	1.00 2.00 1.00 2.00	0
ATOM ATOM	319 320	O N	GLY	37	-5.240	53.828	96.059	1.00 2.00	0
MOTA MOTA	322 323	CA C	GLY GLY	37 37	-6.264 -6.118	52.814 52.251	96.177 97. 57 5	1.00 2.00 1.00 2.00	ő
MOTA	324	ŏ	GLY	37	-7.106	52.030	98.289	1.00 2.00 1.00 8.88	0
MOTA	325 327	N	LEU LEU	38 38	-4.864 -4.558	52.039 51.515	97. 97 5 99.288	1.00 8.88 1.00 8.88	ŏ
MOTA MOTA	327	CA CB	LEU	38	-3.061	51.550	99.559	1.00 2.00	0
MOTA	329	CG	LEU	38 38	-2.202 -0.765	50.559 50.804	98.784 99.171	1.00 2.00 1.00 2.00	Ö
MOTA MOTA	330 331		LEU LEU	38	-2.614	49.106	99.063	1.00 2.00	0
MOTA	332	C	LEU	38 38	-5. 278 -6. 13 0		100.258	1.00 8.88 1.00 2.00	0
ATOM ATOM	333 334	Ŋ	LEU CY5	39	-4.976	53.705	100.216	1.00 2.00	0
ATOM	336	CA	CYS	39	-5.613		101.099	1.00 2.00 1.00 2.00	0
ATOM ATOM	337 338	CB SG	CYS	39 39	-5.339 -3.719	56.686	100.830	1.00 2.00	0
ATOM	339	C	CYS	39	-7.140	54.555	101.201	1.00 2.00 1.00 2.00	0
MOTA	340	0	CYS	39 40	-7.708 -7.793	54.580 54.412	102.296	1.00 2.00 1.00 2.00	0
MOTA MOTA	341 343	N CA	LEU LEU	40	-9.233	54.330	99. 9 99	1.00 2.00	0
ATOM	344	CB	LEU	40	-9 . 73 5	54.704	98.599	1.00 2.00	0

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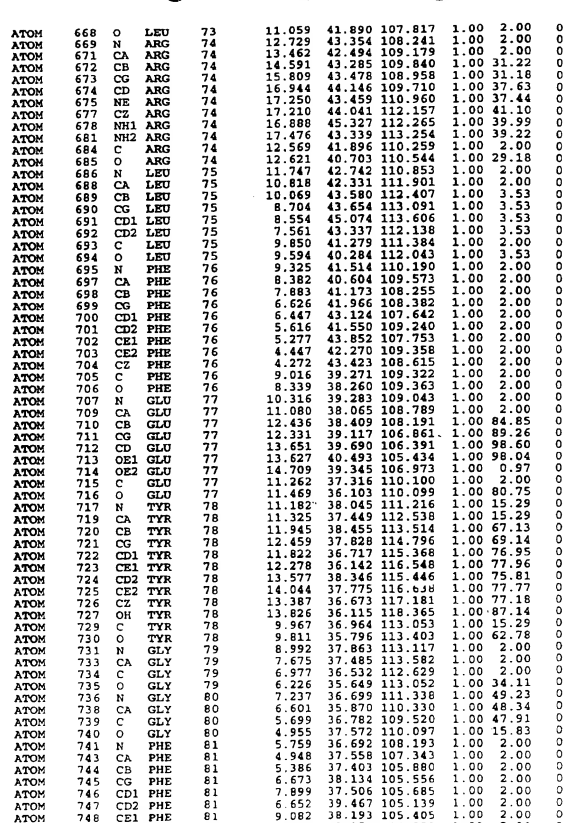
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ATOM	432	OG	SER	48	-14.844	48.045 107.622	1.00 30.63	Ö
ATOM	434	Č	SER	48	-15.624	48.473 110.482	1.00 2.00	0
ATOM	435	Õ	SER	48	-16.184	47.390 110.700	1.00 28.70	0
ATOM	436	N	GLN	49	-14.627	48.922 111.225	1.00 61.72	0
MOTA	438	CA	GLIN	49	-14.175	48.193 112.394	1.00 63.25	0
MOTA	439	CB	GLN	49	-12.763	47.645 112.173	1.00 13.32	0
MOTA	440	CG	GLN	49	-12.668	46.511 111.146	1.00 7.00	0
MOTA	441	CD	GLN	49	-11.246 -10.251	45.952 110.995 46.640 111.258	1.00 8.88 1.00 9.23	0
MOTA	442 443	OE1 NE2	GLN	4 9 4 9	-11.151	44.700 110.573	1.00 7.45	Ö
MOTA MOTA	446	C	GLN	49	-14.210	49.205 113.540	1.00 63.04	ŏ
ATOM	447	ŏ	GLN	49	-13.864	50.373 113.357	1.00 14.61	0
ATOM	448	N	PRO	50	-14.645	48.778 114.734	1.00 34.58	0
ATOM	449	CD	PRO	50	-14.967	47.399 115.118	1.00 4.85	0
MOTA	450	CA	PRO	50	-14.728	49.657 115.901	1.00 34.58	0
ATOM	451	CB	PRO	50	-15.037	48.678 117.032	1.00 2.00	0
MOTA	452	CG	PRO	50	-14.496	47.365 116.531	1.00 2.00 1.00 34.58	0
ATOM	453	Č	PRO	50 50	-13.459 -12.356	50.468 116.164 50.002 115.884	1.00 34.38	0
MOTA	454	0	PRO ILE	50 51	-13.626	51.673 116.708	1.00 2.00	ŏ
MOTA	455 457	N CA	ILE	51	-12.499	52.550 117.015	1.00 2.00	ŏ
MOTA MOTA	458	CB	ILE	51	-13.001	53.975 117.318	1.00 2.00	Ō
ATOM	459	CG2	ILE	51	-13.642	54.035 118.673	1.00 2.00	0
MOTA	460	CG1	ILE	51	-11.850	54.958 117.295	1.00 2.00	0
ATOM	461	CD1	ILE	51	-12.328	56.370 117.351	1.00 2.00	0
MOTA	462	С	ILE	51	-11.650	51.996 118.172	1.00 2.00	0
MOTA	463	0	ILE	51	-10.456	52.303 118.283	1.00 2.00	0
MOTA	464	N	LEU	52	~12.284 ~11.622	51.187 119.028 50.522 120.159	1.00 2.00 1.00 2.00	Ö
ATOM	466	CA	LEU	52 52	-12.391	50.746 121.473	1.00 2.00	ŏ
ATOM ATOM	467 468	CB CG	LEU	52	-11.655	50.518 122.808	1.00 2.00	Ō
ATOM	469		LEU	52	-12.559	50.899 123.944	1.00 2.00	0
ATOM	470		LEU	52	-11.233	49.085 122.973	1.00 2.00	0
ATOM	471	C	LBU	52	-11.660	49.033 119.770	1.00 2.00	0
ATOM	472	0	LEU	52	-12.652	48.331 120.006	1.00 2.00	0
ATOM	473	N	LEU	53	-10.584	48.576 119.136 47.204 118.667	1.00 2.00 1.00 2.00	0
MOTA	475	CA	LEU	53	-10.464 -9.066	46.982 118.069	1.00 2.00	ŏ
ATOM	476	CB	LEU	53 53	-8.802	47.038 116.555	1.00 2.00	ō
MOTA	477 478	CG CD1	LEU	53	-9.835	47.889 115.810	1.00 2.00	0
ATOM ATOM	479		PEO	53	-7.401	47.550 116.353	1.00 2.00	0
ATOM	480	c	LEU	53	-10.686	46.230 119.792	1.00 2.00	0
ATOM	481	õ	LEU	53	-10.365	46.522 120.937	1.00 2.00	0
MOTA	482	N	GLU	54	-11.251	45.076 119.472	1.00 19.77	0
MOTA	484	CA	GLU	54	-11.465	44.049 120.474 43.809 120.714	1.00 20.13 1.00 56.32	0
MOTA	485	CB	GLU	54	-12.955	43.809 120.714 43.362 122.135	1.00 56.32	Ö
MOTA	486	CG	GLU	54	-13.244 -14.668	42.873 122.346	1.00 69.64	Ö
MOTA	487	CD	GLU GLU	54 54	-15.613	43.627 122.024	1.00 78.06	0
MOTA MOTA	488 489		GLU	54	-14.839	41.734 122.848	1.00 73.27	O
MOTA	490	C	GLU	54	-10.798	42.820 119.882	1.00 19.77	Ō
ATOM	491	ŏ	GLU	54	-11.451	41.949 119.313	1.00 48.57	0
ATOM	492	N	LEU	5 5	-9.473	42.800 119.974	1.00 2.00	0
MOTA	494	CA	LEU	55	-8.666	41.712 119.452	1.00 2.00 1.00 2.00	Ö
MOTA	495	CB	LEU	5 5	-7.245 -7.148	42.188 119.176 43.458 118.336	1.00 2.00	Ö
MOTA	496	CG	LEU	55 5 5	-7.148 -5.695	43.728 117.961	1.00 2.00	Ō
ATOM	497		LEU	55	-8.024	43.298 117.108	1.00 2.00	0
MOTA MOTA	498 499	CD2	LEU	5 5	-8.624	40.586 120.456	1.00 2.00	0
MOTA	500	Ö	LEU	5 5	-8.724	40.807 121.664	1.00 10.45	0
MOTA	501	Ň	GLU	56	-8.448	39.374 119.961	1.00 2.00	0
MOTA	503	CA	GLU	5 6	-8.407	38.224 120.826	1.00 2.00	0
MOTA	504	CB	GLU	56	-9.741	37.467 120.729 38.283 121.171	1.00 18.79 1.00 30.34	Ö
MOTA	505	CG	GLU	56	-10.989	38.283 121.171 38.677 122.658	1.00 30.34	ő
ATOM	506	CD	GLU	56 56	-11.012 -11.188	37.789 123.526	1.00 44.70	Ō
ATOM	507	OEI	GLU	, 0	11.100			

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						20 000	100 055	1.00 47.48	0
HOTA	508	OE2		56	-10.872		122.955 120.393	1.00 2.00	Õ
ATOM	509	C	GLU	56	-7.244		119.800	1.00 3.40	ŏ
ATOM	510	0	GLU	56 57	-6.292 -7.312		120.765	1.00 65.90	ŏ
ATOM	511	И	ALA	57 57	-6.336		120.405	1.00 62.88	ŏ
ATOM	513	CA	ALA ALA	57	-6.750	34.434	119.039	1.00 26.61	ŏ
MOTA	514 515	CB C	ALA	57 57	-4.867		120.387	1.00 68.03	Õ
ATOM	516	0	ALA ALA	57	-4.543		120.932	1.00 14.05	ŏ
MOTA MOTA	517	N	PRO	58	-3.951		119.857	1.00 2.00	Ō
ATOM	518	CD	PRO	58	-4.109		119.533	1.00 9.23	0
ATOM	519	CA	PRO	58	-2.526	34.988	119.782	1.00 2.00	0
ATOM	520	CB	PRO	58	-1.852		119.501	1.00 9.23	0
ATOM	521	CG	PRO	58	-2.881		118.711	1.00 9.49	0
MOTA	522	C	PRO	58	-2.176		118.672	1.00 2.00	0
ATOM	523	0	PRO	58	-2.688		117.540	1.00 12.29	0
MOTA	524	N	LEU	59	-1.284		118.975	1.00 17.05	0
ATOM	526	CA	LEU	59	-0.856		117.989	1.00 17.05	0
MOTA	527	CB	LEU	59	-1.862		117.895	1.00 2.00	0
MOTA	528	CC	LEU	59	-1.842		118.993	1.00 2.00	0
MOTA	529		LEU	59	-2.880		118.706	1.00 2.00 1.00 2.00	0
MOTA	530		LEU	59 50	-2.117		120.339 118.351	1.00 17.05	ő
ATOM	531	C	LEU	59 50	0.510 1.030		119.429	1.00 2.00	ŏ
ATOM	532	0	LEU	59 60	1.103		117.438	1.00 21.64	ŏ
ATOM	533	N	LYS	60 60	2.399		117.688	1.00 21.64	ŏ
ATOM	53 5	CA	LYS LYS	60	3.358		116.513	1.00 10.44	ŏ
MOTA	536 537	CB CG	LYS	60	3.185		115.768	1.00 11.70	ŏ
MOTA	53 <i>1</i> 538	CD	LYS	60	3.556		116.608	1.00 12.19	Õ
MOTA MOTA	539	CE	LYS	60	5.034		116.544	1.00 8.37	0
ATOM	540	NZ	LYS	60	5.824		117.066	1.00 8.37	0
ATOM	544	C	LYS	60	2.063		117.804	1.00 21.64	0
ATOM	545	ŏ	LYS	60	1.142	41.832	117.128	1.00 13.47	0
MOTA	546	N	ILE	61	2.757	42.072	118.680	1.00 20.60	0
ATOM	548	CA	ILE	61	2. 49 9		118.822	1.00 22.32	0
ATOM	549	CB	ILE	61	2.032	43.859	120.240	1.00 2.00	0
MOTA	550	CG2	ILE	61	1.485	45.288	120.239	1.00 2.00	0
MOTA MOTA	550 551	CG2 CG1	ILE	61	1.485 0.940	45.288 42.881	120.239 120.702	1.00 2.00	0
	551 552	CG1 CD1	ILE ILE ILE	61 61	1.485 0.940 0.019	45.288 42.881 43.422	120.239 120.702 121.783	1.00 2.00 1.00 2.00	0
MOTA MOTA MOTA	551 552 553	CG1 CD1 C	ILE ILE ILE	61 61 61	1.485 0.940 0.019 3.791	45.288 42.881 43.422 44.220	120.239 120.702 121.783 118.494	1.00 2.00 1.00 2.00 1.00 23.52	0 0
MOTA MOTA MOTA	551 552 553 554	CG1 CD1 C	ILE ILE ILE ILE	61 61 61 61	1.485 0.940 0.019 3.791 4.862	45.288 42.881 43.422 44.220 43.752	120.239 120.702 121.783 118.494 118.868	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00	0 0 0
ATOM ATOM ATOM ATOM	551 552 553 554 555	CG1 CD1 C O N	ILE ILE ILE ILE CYS	61 61 61 61 62	1.485 0.940 0.019 3.791 4.862 3.698	45.288 42.881 43.422 44.220 43.752 45.341	120.239 120.702 121.783 118.494 118.868 117.787	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00	0 0 0 0
ATOM ATOM ATOM ATOM ATOM	551 552 553 554 555 557	CG1 CD1 C O N CA	ILE ILE ILE ILE CYS CYS	61 61 61 62 62	1.485 0.940 0.019 3.791 4.862 3.698 4.874	45.288 42.881 43.422 44.220 43.752 45.341 46.095	120.239 120.702 121.783 118.494 118.868 117.787 117.393	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00	0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 552 553 554 555 557 558	CG1 CD1 C O N CA CB	ILE ILE ILE ILE CYS CYS CYS	61 61 61 62 62 62	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47	0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 552 553 554 555 557 558 559	CG1 CD1 C O N CA CB SG	ILE ILE ILE ILE CYS CYS CYS	61 61 61 62 62 62 62	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 44.252	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47	0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 552 553 554 555 557 558 559 560	CG1 CD1 C O N CA CB SG C	ILE ILE ILE ILE CYS CYS CYS CYS	61 61 61 62 62 62 62 62	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 44.252 47.560	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47	0000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 552 553 554 555 557 558 559 560 561	CG1 CD1 C O N CA CB SG C	ILE ILE ILE ILE CYS CYS CYS CYS CYS	61 61 61 62 62 62 62 62 62	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 44.252 47.560 47.957	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 1.00 1.00 11.47 1.00 11.47	00000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	551 552 553 554 555 557 558 560 561 562	CG1 CD1 C O N CA CB SG C O N	ILE ILE ILE ILE CYS CYS CYS CYS CYS CYS	61 61 61 62 62 62 62 62 62 63	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 44.252 47.560 47.957	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658	1.00 2.00 1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 1.00 1.00 11.47 1.00 2.00 1.00 11.47	0000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	551 552 553 554 555 557 558 559 560 561	CG1 CD1 C O N CA CB SG C	ILE ILE ILE ILE CYS CYS CYS CYS CYS	61 61 61 62 62 62 62 62 62	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 44.252 47.560 47.957 48.367 49.610	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5555 5555 5561 5566 5665 5665	CG1 C O N CA CB SG C O N CA C	ILE ILE ILE ILE CYS CYS CYS CYS CYS CYS GLY GLY	61 61 61 62 62 62 62 62 63 63	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 44.252 47.560 47.957 48.367 49.789 50.610 50.109	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.971 117.222 116.897	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.39	00000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	551 552 553 554 555 557 558 559 560 561 562 564	CG1 CD1 CONCA CBSG CONCA	ILE ILE ILE ILE CYS CYS CYS CYS CYS CYS CYS	61 61 61 62 62 62 62 62 63 63	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548	45.288 42.881 43.422 44.220 43.752 45.925 45.925 44.252 47.560 47.957 48.367 49.789 50.610 50.109 51.870	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.971 117.222 116.897 116.959	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 2.00	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5557 5557 5560 5664 5664 5666 5667 569	CG1 CD1 CONCACB SGCONCACO	ILE ILE ILE ILE CYS CYS CYS CYS CYS GLY GLY GLY GLY ASP	61 61 61 62 62 62 62 63 63 63 64	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548 7.624 6.216 7.068	45.288 42.881 43.422 44.220 43.752 45.925 45.925 44.252 47.560 47.957 48.367 49.789 50.610 50.610 51.870 52.834	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 2.00 1.00 11.47 1.00 2.00 1.00 1.39 1.00 2.00 1.00 3.88	0000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5557 5557 5560 5662 5665 56667 5667 567	CG1 CD1 CONCA CBSG CONCA CONCA CCB	ILE ILE ILE ILE CYS CYS CYS CYS GLY GLY GLY ASP ASP	61 61 61 62 62 62 62 63 63 63 64 64	1.485 0.940 0.019 3.791 4.862 3.6874 5.147 5.439 4.607 3.451 5.662 5.507 6.548 7.624 67.068 7.805	45.288 42.881 43.422 44.220 43.752 45.341 46.095 44.252 47.560 47.957 48.367 49.789 50.610 50.109 51.870 52.834 53.721	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 3.88 1.00 6.62	00000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5557 5557 5559 5661 5664 56667 570 571	CG1 CD1 CONCA CBSC CONCA CONCA CBCC	ILE ILE ILE ILE CYS CYS CYS CYS GLY GLY GLY ASP ASP	61 61 61 62 62 62 62 63 63 64 64 64	1.485 0.940 0.019 3.791 4.862 3.6874 5.439 4.607 3.451 5.662 5.507 6.548 7.624 67.068 7.805 6.873	45.288 42.881 43.422 44.220 43.752 45.341 46.095 45.925 47.560 47.957 48.367 49.789 50.610 50.109 51.870 52.834 53.721 54.589	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245 118.025	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 11.39 1.00 2.00 1.00 3.88 1.00 6.62 1.00 6.62	000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5557 5557 5559 5661 5667 5679 5712	CG1 CD1 CONCACBSCONCACONCACCONCACCOONCACOONCACCOONC	ILE ILE ILE ILE CYS CYS CYS CYS CYS GLY GLY GLY ASP ASP ASP	61 61 62 62 62 62 63 63 64 64 64	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548 7.624 6.216 7.805 6.873 5.996	45.288 42.881 43.422 44.220 43.752 45.925 45.925 47.560 47.957 48.367 49.789 50.610 50.109 51.870 52.834 53.721 54.019	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.739 117.222 116.897 116.959 116.255 117.245 118.025 118.708	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 11.39 1.00 2.00 1.00 1.388 1.00 6.62 1.00 6.62 1.00 12.70	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5557 5557 5559 5661 5667 567 5772 573	CG1 CD1 CONCACB SCONCACCONCACCO	ILE ILE ILE ILE ILE CYS CYS CYS CYS CYS GLY GLY GLY ASP ASP ASP	61 61 61 62 62 62 62 63 63 64 64 64	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548 7.624 6.216 7.068 7.805 6.873 5.996 7.003	45.288 42.881 43.422 44.220 43.752 45.925 45.925 44.252 47.560 47.957 48.367 49.789 50.610 50.109 51.870 52.834 53.721 54.589 55.829	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245 118.025 118.708 117.946	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 2.00 1.00 11.47 1.00 2.00 1.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 3.88 1.00 6.62 1.00 6.62 1.00 12.70	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5555 5555 5566 5667 5667 5771 5773 574	CG1 CD1 CONCACB CCONCACC CONCACC COD1 CCB CCOD1 CCB CCOCC	ILE ILE ILE ILE CYS CYS CYS CYS GLY GLY GLY ASP ASP ASP ASP ASP	61 61 61 62 62 62 62 63 63 64 64 64 64	1.485 0.940 0.019 3.791 4.862 3.698 4.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548 7.624 6.216 7.068 7.805 6.873 5.903 8.064	45.288 42.881 43.422 44.220 43.752 45.925 44.252 47.560 47.957 48.367 49.789 50.610 50.610 51.870 52.834 53.721 54.589 54.019 55.829 52.362	120.239 120.702 121.783 118.494 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.255 117.245 118.025 118.708 117.946 115.221	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 3.88 1.00 6.62 1.00 6.62 1.00 12.70 1.00 12.70 1.00 14.37	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5555 5555 5566 55667 5577 5777 57	CG1 CD1 CONCA CBSG CONCA CBCG OD1 CACCOONCA CBOD1	ILE ILE ILE ILE CYS CYS CYS CYS GLY GLY GLY ASP ASP ASP ASP ASP	61 61 61 62 62 62 62 63 63 64 64 64 64 64	1.485 0.940 0.019 3.791 4.862 3.874 5.147 5.439 4.607 3.451 5.662 5.548 7.624 6.216 7.068 7.805 6.873 5.903 8.064 9.221	45.288 42.881 43.422 44.220 43.752 45.925 46.095 45.925 47.560 47.957 48.367 49.789 50.610 50.109 51.870 52.834 53.721 54.589 54.019 55.829 54.019 55.829 52.362 52.080	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.255 117.245 118.025 118.708 117.946 115.221 115.557	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 1.00 1.39 1.00 2.00 1.00 3.88 1.00 6.62 1.00 6.62 1.00 12.70 1.00 12.70 1.00 14.37 1.00 8.57	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5555 5555 5566 55667 5577 5777 57	CG1 CD1 CONCA CBSG CONCA CCBCDD2 CONCA CBCDD2 CONCA	ILE ILE ILE ILE ILE CYS CYS CYS CYS GLY GLY GLY ASP ASP ASP ASP ASP ASP	61 61 61 62 62 62 62 63 63 64 64 64 64 64 65	1.485 0.940 0.019 3.791 4.862 3.874 5.147 5.439 4.607 3.451 5.662 5.507 6.548 7.805 6.216 7.068 7.805 6.873 5.903 8.064 9.221 7.619	45.288 42.881 43.422 44.220 43.752 45.395 46.095 46.925 47.560 47.957 48.367 49.789 50.6109 50.870 52.834 53.721 54.589 54.019 55.829 52.360 52.360 52.360	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245 118.025 118.708 117.946 115.221 115.557 113.963	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 11.39 1.00 2.00 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 13.71 1.00 8.57 1.00 8.57	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5557 5556 55664 55667 5777 5777 57	CG1 CD1 CONCA CBSG CONCA CBCGD1 CCD0 CONCA	ILE	61 61 61 62 62 62 62 63 63 64 64 64 64 65 65	1.485 0.940 0.019 3.791 4.862 3.6874 5.439 4.607 3.451 5.662 5.504 7.6216 7.068 7.805 6.873 5.996 8.064 9.219 8.471	45.288 42.881 43.422 44.220 43.752 45.395 44.252 47.560 47.957 48.367 49.789 50.6109 50.870 52.834 53.721 54.589 54.019 55.829 52.360 52.360 51.924	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.255 117.245 118.025 118.708 117.946 115.221 115.557 113.963 112.849	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 11.39 1.00 2.00 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 13.71 1.00 8.57 1.00 8.57 1.00 5.19 1.00 3.07	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5555 5555 5566 55666 55666 5577 5777 5777 5778 5779	CG1 CD1 CONCACOONCACOODOCONCACOONCACOONCACOO	ILE	61 61 61 62 62 62 62 63 63 64 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.862 3.6874 5.439 4.607 3.451 5.662 5.507 6.548 7.6216 7.068 7.805 6.873 5.996 7.0068 7.0	45.288 42.881 43.422 44.220 43.752 45.341 46.925 47.560 47.957 48.367 49.789 50.6109 50.870 52.834 53.721 54.589 54.019 55.829 52.360 52.360 53.360 5	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.266 117.658 117.751 117.739 117.971 117.921 116.897 116.897 116.255 118.025 118.025 118.025 118.025 118.557 115.557 115.557 115.563 112.849 111.672	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 11.39 1.00 3.88 1.00 6.62 1.00 12.70 1.00 12.70 1.00 12.70 1.00 13.71 1.00 8.57 1.00 3.07 1.00 3.07	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5555 5555 5555 5555 5566 55667 5577 5777 5777 5777 5779 5779	CG1 CD1 CONCACONCACOONCACBCCOONCACBCCCCCCCCCCCCC	ILE	61 61 61 62 62 62 62 63 63 64 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.8698 4.874 5.147 5.439 4.607 3.451 5.5607 6.548 7.624 6.216 7.805 8.993 8.064 9.221 7.619 8.4609	45.288 42.881 43.422 44.220 43.752 45.925 45.925 47.560 47.957 48.369 50.610 50.109 51.870 52.834 53.721 54.519 55.829 52.326 52.326 51.924 50.903	120.239 120.702 121.783 118.494 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245 118.025 118.708 117.946 115.221 115.557 113.963 112.849 111.849 111.855	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 11.39 1.00 3.88 1.00 6.62 1.00 12.70 1.00 12.70 1.00 12.70 1.00 13.71 1.00 8.57 1.00 3.07 1.00 3.07	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	551 5553 5555 5555 5555 5566 55667 5577 5777 57	CG1 CD1 CONCA CBSC CONCA CBCCOD1 CBCCCONCA CBCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ILE	61 61 61 62 62 62 62 63 63 63 64 64 64 64 64 64 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.862 3.6874 5.439 4.607 3.451 5.662 5.507 6.548 7.6216 7.068 7.805 6.873 5.996 7.0068 7.0	45.288 42.881 43.422 44.220 43.752 45.925 44.252 47.560 47.957 48.367 49.567 49	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.266 117.658 117.751 117.739 117.971 117.921 116.897 116.897 116.255 118.025 118.025 118.025 118.025 118.557 115.557 115.557 115.563 112.849 111.672	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 11.39 1.00 6.62 1.00 6.62 1.00 6.62 1.00 12.70 1.00 12.70 1.00 13.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 12.70 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	55555555555555555555555555555555555555	CG1 CD1 CONCACONCACOONCACCOONCACCCCCCCCCCCCCCCC	ILE	61 61 61 61 62 62 62 63 63 63 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.8698 4.874 5.147 5.439 4.607 3.451 5.6607 6.548 7.624 6.216 7.805 8.996 7.903 8.064 9.221 7.669 8.471 8.471 7.669 6.871	45.288 42.881 43.422 44.220 43.752 45.925 44.252 47.560 47.957 48.369 50.610 50.109 51.870 52.834 53.721 54.589 52.362 52.362 52.326 51.924 51.310 50.095 49.023	120.239 120.702 121.783 118.494 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245 118.025	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 11.47 1.00 2.00 1.00 11.39 1.00 2.00 1.00 11.39 1.00 6.62 1.00 6.62 1.00 12.70 1.00 12.70 1.00 14.37 1.00 8.57 1.00 8.57 1.00 5.19 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	55555555555555555555555555555555555555	CG1 CD1 CONCA CBSG CONCA CCONCA CCGOOD2 CONCA CCGCOONCA CCGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ILE	61 61 61 62 62 62 62 63 63 63 64 64 64 64 64 64 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.8698 4.874 5.147 5.439 4.607 3.451 5.662 5.548 7.624 6.216 7.805 8.873 5.903 8.064 9.221 7.663 8.673 7.663 8.671 7.663 8.671 7.723	45.288 42.881 43.422 44.220 43.752 45.925 44.252 47.560 47.957 48.367 49.789 50.610 50.610 51.870 52.834 53.721 54.589 54.019 55.829 54.019 52.362 51.924 51.310 50.905 51.870	120.239 120.702 121.783 118.494 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.222 116.897 116.959 116.255 117.245 118.025	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 2.00 1.00 11.39 1.00 2.00 1.00 3.88 1.00 6.62 1.00 12.70 1.00 12.70 1.00 12.70 1.00 14.37 1.00 8.57 1.00 3.07 1.00 2.00 1.00 2.00 1.00 2.00 1.00 3.07 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	55555555555555555555555555555555555555	CG1 CD1 CONCA CBSG CONCA CBCOD1 CONCA CBCCD1 CCD1 CCD1 CCD1	ILE	61 61 61 61 62 62 62 63 63 63 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.862 3.874 5.147 5.439 4.607 3.451 5.662 5.548 7.805 6.873 5.903 8.064 9.221 7.663 8.609 8.471 7.663 8.609 8.71 7.723 9.185	45.288 42.881 43.422 44.222 45.752 45.925 46.925 47.560 47.957 48.789 50.6109 51.870 52.834 53.789 54.019 55.821 54.589 54.019 55.820 52.326 51.320 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 5	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.971 117.921 116.897 116.255 117.245 118.025 118.708 117.946 115.557 113.963 112.849 111.672 110.535 112.151 112.357 112.357	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 2.00 1.00 1.39 1.00 2.00 1.00 1.39 1.00 2.00 1.00 12.70 1.00 12.70 1.00 12.70 1.00 14.37 1.00 8.57 1.00 5.19 1.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	55555555555555555555555555555555555555	CG1 CD1 CONCA CBSG CONCA CCONCA CCGOOD2 CONCA CCGCOONCA CCGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ILE	61 61 61 61 61 61 62 62 62 63 63 63 64 64 64 64 64 64 65 65 65 65 65 65 65 65 65 65 65 65 65	1.485 0.940 0.019 3.791 4.862 3.874 5.147 5.439 4.607 3.451 5.662 5.548 7.805 6.873 5.903 8.064 9.221 7.663 8.609 6.871 7.663 8.609 6.71 7.723 9.185 10.379	45.288 42.881 43.422 44.222 45.752 45.925 46.925 47.560 47.957 48.789 50.6109 51.870 52.834 53.789 54.019 55.821 54.589 54.019 55.820 52.326 51.320 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 50.925 51.310 5	120.239 120.702 121.783 118.494 118.868 117.787 117.393 115.895 115.266 117.658 117.751 117.739 117.222 116.897 116.255 118.025 118.025 118.708 117.946 115.221 115.557 113.963 112.849 111.672 110.535 112.151 112.763 112.357 112.101	1.00 2.00 1.00 23.52 1.00 2.00 1.00 2.00 1.00 2.00 1.00 11.47 1.00 11.47 1.00 2.00 1.00 1.00 1.00 2.00 1.00 2.00 1.00 1.39 1.00 2.00 1.00 3.88 1.00 6.62 1.00 6.62 1.00 12.70 1.00 12.70 1.00 14.37 1.00 8.57 1.00 5.19 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000

ATOM ATOM	588 589	C O	HIS HIS	66 66	9.847 11.042	55.525 110.502 55.818 110.539	1.00 2.00 1.00 12.38	0
ATOM	590	CB	HIS	66	9.846	56.182 112.881	1.00 8.25 1.00 8.25	0
ATOM	591	CG.	HIS	66	9.040 8.104	57.046 113.782 57.949 113.337	1.00 8.25	ŏ
MOTA	592 594		HIS HIS	66 66	9.001	57.109 115.133	1.00 8.25	ŏ
ATOM ATOM	595		HIS	66	8.042	58.043 115.526	1.00 8.25	0
ATOM	596		HIS	66	7.536	58.516 114.399	1.00 8.25	0
ATOM	597	N	GLY	67	9.197	55.195 109.385	1.00 11.52 1.00 11.52	0
MOTA	599	CA	GLY	67	9.835 11.029	55.174 108.078 54.266 107.868	1.00 11.52 1.00 11.52	0
ATOM	600	C	GLY GLY	67 67	11.782	54.462 106.915	1.00 81.94	ŏ
MOTA MOTA	601 602	O N	GLN	68	11.214	53.272 108.730	1.00 18.96	0
ATOM	604	CA	GLN	68	12.353	52.364 108.602	1.00 20.32	0
MOTA	605	CB	GTN	68	13.007	52.139 109.957	1.00 6.39	0
MOTA	606	CG	GLN	68	13.261 14.315	53.407 110.711 53.239 111.757	1.00 4.06 1.00 7.01	0
MOTA	607	CD	GLN	68 68	15.013	54.197 112.111	1.00 8.21	ŏ
MOTA	608 609	OE1 NE2	GLN GLN	68	14.453	52.020 112.270	1.00 5.84	Ō
MOTA MOTA	612	C	GLN	68	11.916	51.039 108.010	1.00 18.72	0
ATOM	613	ŏ	GLN	68	12.01B	49.975 108.634	1.00 10.73	0
MOTA	614	N	TYR	69	11.450	51.119 106.777 49.970 106.054	1.00 5.70 1.00 5.52	0
MOTA	616	CA	TYR	69 60	10.959 10.807	50.313 104.587	1.00 2.00	ŏ
MOTA	617	CB CC	TYR TYR	69 69	9.988	49.322 103.841	1.00 2.00	Ō
MOTA MOTA	618 619	CG CD1	TYR	69	8.658	49.099 104.187	1.00 2.00	0
ATOM	620	CEI	TYR	69	7.873	48.194 103.477	1.00 2.00	0
ATOM	621	CD2	TYR	69	10.526	48.611 102.762	1.00 2.00 1.00 2.00	0
MOTA	622	CE2	TYR	69	9.751 8. 4 31	47.701 102.038 47.506 102.408	1.00 2.00	Ö
ATOM	623	CZ	TYR	69 69	7.656	46.632 101.715	1.00 2.00	ō
ATOM ATOM	624 626	OH C	TYR TYR	69	. 11.815	48.730 106.188	1.00 8.52	0
ATOM	627	Ö	TYR	69	11.284	47.641 106.393	1.00 2.00	0
ATOM	628	N	TYR	70	13.132	48.877 106.093	1.00 2.00 1.00 2.00	0
MOTA	630	CA	TYR	70 70	13.992 15.420	47.703 106.186 48.031 105.741	1.00 64.80	ŏ
MOTA	631	CB	TYR TYR	70 70	15.533	47.964 104.231	1.00 70.91	0
ATOM ATOM	632 633	CG CD1		70 70	15.026	48.993 103.431	1.00 73.06	0
ATOM	634	CE1	TYR	70	15.082	48.924 102.041	1.00 68.38 1.00 73.86	0
ATOM	635	CD2	TYR	70	16.107	46.857 103.597 46.780 102.200	1.00 73.86 1.00 70.94	ŏ
MOTA	636	_	TYR	70 70	16.171 15.654	47.820 101.433	1.00 72.52	Ō
ATOM	637	CZ OH	TYR TYR	70 70	15.712	47.779 100.062	1.00 70.35	0
ATOM ATOM	638 640	C	TYR	70	13.943	47.007 107.538	1.00 2.00	0
ATOM	641	ŏ	TYR	70	13.967	45.771 107.610	1.00 64.08 1.00 7.87	0
ATOM	642	N	ASP	71	13.821	47.789 108.608 47.203 109.927	1.00 7.87 1.00 6.73	ŏ
MOTA	644	CA	ASP	71	13.728 14.030	48.242 110.998	1.00 14.88	Ō
MOTA	645	CB CG	ASP ASP	71 71	15.514	48.600 111.054	1.00 25.87	0
MOTA MOTA	646 647		ASP	71	15.826	49.778 111.293	1.00 24.58	0
ATOM	648		ASP	71	16.375	47.714 110.861	1.00 21.52 1.00 6.73	0
MOTA	649	C	ASP	71	12.331 12.116	46.622 110.051 45.697 110.826	1.00 14.14	ŏ
MOTA	650	0	ASP	71 72	11.399	47.148 109.250	1.00 2.00	0
MOTA MOTA	651 653	N CA	LEU	72	10.015	46.654 109.208	1.00 2.00	0
MOTA	654	CB	LEU	72	9.094	47.613 108.456	1.00 2.00 1.00 2.00	0
ATOM	655	CG	LEU	72	7.771	46.946 108.067	1.00 2.00 1.00 2.00	ő
MOTA	656		LEU	72	7.025	46.562 109.342 47.867 107.208	1.00 2.00	ŏ
ATOM	657		LEU	72 7 2	6.935 10.014	45,320 108,476	1.00 2.00	0
MOTA	658	C	L E U L E U	72	9.259	44.401 108.814	1.00 2.00	0
MOTA MOTA	659 660	O N	LEU	73	10.848	45.231 107.449	1.00 44.35	0
MOTA	662	CA	LEU	7 3	10.968	44.005 106.693	1.00 42.59 1.00 2.00	0
ATOM	663	CB	LEU	73	11.846 11.248	44.218 105.460 44.759 104.160	1.00 2.00	ő
MOTA	664	CG	LEU	7 3	12.324	44.728 103.101	1.00 2.00	0
MOTA	665	CD	LEU LEU	73 73	10.052	43.920 103.725	1.00 2.00	0
MOTA	666		LEU	73	11.603	42.978 107.629	1.00 42.42	0
MOTA	667	С	1,500	, ,				

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CE2 PHE

749

ATOM

81

7.829

40.154 104.859

1.00

2.00

		0.1	9.044 39.517 104.992 1.00 2.00 0
MOTA	750 CZ PHE	81 81	3.428 37.380 107.548 1.00 2.00 0
ATOM	751 C PHE	81	2.918 36.257 107.636 1.00 2.00 0
ATOM	752 O PHE 753 N PRO	82	2.694 38.505 107.618 1.00 2.00 0
MOTA	753 N PRO 754 CD PRO	82	3 317 39 807 107 337 1.00 21.23 0
MOTA	755 CA PRO	82	1.261 38.702 107.820 1.00 2.00 0
MOTA	756 CB PRO	82	0.996 39.979 107.079 1.00 20.78 0
MOTA MOTA	757 CG PRO	82	2.144 40.765 107.502 1.00 22.98 0
ATOM	758 C PRO	82	0.273 37.623 107.490 1.00 2.00 0
MOTA	759 O PRO	82	-0.761 37.529 108.161 1.00 36.87 0 0.501 36.841 106.423 1.00 27.66 0
ATOM	760 N PRO	83	0.301 30.042 400.000
ATOM	761 CD PRO	83	1.433
MOTA	762 CA PRO	83	-0.514 35.812 106.201 1.00 30.55 0 0.089 34.958 105.083 1.00 5.44 0
MOTA	763 CB PRO	83	1.550 35.427 104.971 1.00 10.08 0
MOTA	764 CG PRO	83	-0.791 34.990 107.460 1.00 24.53 0
MOTA	765 C PRO	83 83	1 947 34 863 107 882 1.00 5.06 0
MOTA	766 O PRO 767 N GLU	84	0.284 34.504 108.080 1.00 63.06 0
MOTA	767 N GLU 769 CA GLU	84	0.190 33.665 109.268 1.00 65.66 0
MOTA	770 CB GLU	84	1.448 32.797 109.382 1.00 61.87 0
MOTA MOTA	771 CG GLU	84	1.194 31.402 109.979 1.00 75.07 0
ATOM	772 CD GLU	84	0.401 30.432 103.00
MOTA	773 OE1 GLU	84	1.015 29.500 100.500
ATOM	774 OE2 GLU	84	110 500 1 00 66 47 0
MOTA	775 C GLU	84	-0.066 34.394 110.588 1.00 60.47 -1.194 34.406 111.085 1.00 61.22 0
MOTA	776 O GLU	84	0.984 34.978 111.161 1.00 66.52 0
ATOM	777 N SER	85 85	0 882 35 691 112.431 1.00 64.76
MOTA	779 CA SER	85	2.266 36.145 112.900 1.00 2.00 0
ATOM	780 CB SER 781 OG SER	85	3.198 35.078 112.871 1.00 2.00 0
MOTA	783 C SER	85	-0.013 36.917 112.349 1.00 62.83 0
MOTA MOTA	784 O SER	85	_0.005 37.633 111.311
MOTA	785 N ASN	86	-0.785 37.140 113.422 1 00 2 00
ATOM	787 CA ASN	86	-1.04/ 30.321 113.43
MOTA	788 CB ASN	86	-2./4/ 38.123 114.304 1 00 10 84 0
MOTA	789 CG ASN	86	-3.887 37.283 113.994 1.00 10.04 -3.763 36.629 112.957 1.00 12.15 0
MOTA	790 OD1 ASN	86 86	-5 017 37 300 114 705 1.00 6.45 0
MOTA	791 ND2 ASN 794 C ASN	86	-0 742 39.467 113.912 1.00 2.00 0
MOTA	794 C ASN 795 O ASN	86	0.363 39.229 114.423 1.00 2.00 0
MOTA MOTA	796 N TYR	87	-1.183 40.703 113.723 1.00 3.00
ATOM	798 CA TYR	87	-0.337 41.830 114.000 1.00 2.00
ATOM	799 CB TYR	87	0.529 \$2.233 112.604 1.00 3.00
ATOM	800 CG TYR	87	1.802 41.446 112.001
MOTA	801 CD1 TYR	87	1.879 40.421 111.749 1.00 2.00 0 3.039 39.689 111.598 1.00 2.00 0
MOTA	802 CE1 TYR	87	2 930 41 714 113 458 1.00 2.00
MOTA	803 CD2 TYR	87 87	4 080 40.991 113.313 1.00 2.00 0
MOTA	804 CE2 TYR 805 CZ TYR	87	4 127 39 986 112.384 1.00 2.00
MOTA	805 CZ TYR 806 OH TYR	87	5 280 39 281 112,259 1.00 2.00 U
MOTA MOTA	808 C TYR	87	-1.104 43.029 114.588 1.00 2.00 0
MOTA	809 O TYR	87	-2.264 43.238 114.216 1.00 2.00 0
MOTA	810 N LEU	88	-0.440 43.810 113.433 1.00 2.00 0
MOTA	812 CA LEU	88	-1.015 45.026 113.307 1.00 2.00 0
MOTA	813 CB LEU	88	3 364 45 917 118.062 1.00 2.00 0
MOTA	814 CG LEU	88	_3 657 46.225 117.348 1.00 2.00 0
MOTA	815 CD1 LEU	88 88	-2 675 45.551 119.493 1.00 2.00 0
MOTA	816 CD2 LEU 817 C LEU	88	0 119 46.034 116.084 1.00 2.00
MOTA		88	1 082 45.814 116.814 1.00 2.00 0
MOTA	818 O LEU 819 N PHE	89	0 061 47.108 115.309 1.00 2.00 0
ATOM	821 CA PHE	89	1.114 48.117 115.393 1.00 2.00 0
MOTA MOTA	822 CB PHE	89	1.567 48.599 114.004 1100 2 00 0
MOTA	823 CG PHE	89	2.303
ATOM	824 CD1 PHE	89	1.61/ 46.543 112.150 1.00 2.00 0
ATOM	825 CD2 PHE	89	3.683 47.567 111.885 1.00 2.00 0
MOTA	826 CE1 PHE	89	4.379 46.586 112.464 1.00 2.00 0
MOTA	827 CE2 PHE	89	41310 - 4222

ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	829 833 8334 8334 8336 8336 8336 8442 8443 8445 8445 8449 849	C O N CA C O N CA C C C	PHE PHE LEU LEU LEU LEU LEU LEU GLY GLY GLY ASP ASP ASP	89 89 90 90 90 91 91 91 92 92 92 92	3.673 0.495 -0.664 1.271 0.810 1.519 1.516 2.488 0.111 1.000 1.382 0.734 0.884 2.307 3.220 2.471 3.736 4.185 3.449	50.856 50.702 49.291 49.246 48.870 52.275 53.196 52.459 53.774 54.139 53.312 55.400 55.978 56.170 57.239 58.145	116.197 115.985 117.124 118.014 119.991 121.147 120.402 117.463 118.200 116.175 115.588 115.209 115.267 114.820 114.382 115.576 116.525 116.063	1.00 1.00 1.00 1.00 1.00 1.00	14.03 14.03 32.27 2.00 2.00 2.00 2.00 2.00	000000000000000000000000000000000000000
MOTA	851	OD2	ASP	92	4.541	57.177	117.725 113.259	1.00	2.00	0
MOTA MOTA	852 853	C O	ASP ASP	92 92	5.515	54.631	113.422	1.00	2.00	0
HOTA	854	N	TYR	93 93	3.792 4.301		112.096 110.938	1.00	2.00	0
MOTA MOTA	856 857	CA CB	TYR TYR	93	3.149	53.907	110.171	1.00	2.00	0
ATOM	858	CG	TYR	93	2.122 0.875		111.029 111.227	1.00	2.00	0
MOTA MOTA	859 860	CD1 CE1		93 93	-0.086	53.176	112.004	1.00	2.00	0
ATOM	861	CD2	TYR	93	2.390		111.628 112.405	1.00	2.00	0
MOTA	862	CE2 CZ	TYR TYR	93 93	1.445 0.204	51.951	112.593	1.00	2.00	0
ATOM ATOM	863 864	OH	TYR	93	-0.736		113.379	1.00	2.00	0
MOTA	866	C	TYR	93 93	5.088 5.998		109.992 109.335	1.00	2.00	Ö
atom atom	867 868	N O	TYR VAL	94	4.718	56.667	109.904	1.00	2.00	0
ATOM	870	CA	VAL	94	5.380 4.322		109.004 108.112	1.00 1.00	2.00	0
MOTA MOTA	871 872	CB CG1	VAL	94 94	3.365	57.315	107.523	1.00	2.00	0
ATOM	873	CG2		94	3.551		108.903	1.00 1.00	2.00	0
MOTA	874	C	VAL	94 94	6.253. 6.447	58.614 58.457	109.780 110.985	1.00	2.00	Ö
MOTA MOTA	875 876	O N	VAL ASP	95	6.774	59.629	109.094	1.00	8.40	0
MOTA	878	CA	ASP	95	7.641		109.689 110.868	1.00	2.00	0
MOTA	879	CB CG	ASP ASP	95 95	6.967 5.975	62.433	110.449	1.00	45.35	0
ATOM ATOM	880 881		ASP	95	6.227	63.167	109.471	1.00	44.75 53.79	0 0
MOTA	882	OD2		95 05	4.937 8.975		111.127 110.155	1.00	2.71	ŏ
MOTA MOTA	883 884	C	ASP ASP	95 95	9.092	58.877	110.421	1.00	37.69	0
MOTA	885	N	ARG	96	9.972	60.949	110.252 110.685	1.00	21.97 27.39	0
MOTA	887 888	CA CB	ARG ARG	96 96	11.322 11.285	59.989	112.099	1.00	18.63	0
MOTA MOTA	889	CG	ARG	96	12.037		113.128	1.00	24.81 32.64	0
MOTA	890	CD	ARG	96 06	11.255 10.551	62.052 61.833	113.581 114.850	1.00	39.46	ŏ
MOTA MOTA	891 893	NE CZ	ARG ARG	96 96	9.890	62.774	115.525	1.00	39.86	0
MOTA	894	NH	ARG	96	9.824	64.018	115.066 116.670		40.95	0 0
MOTA	897	NH2		96 96	9.290 12.109	59.659	109.732	1.00	24.24	0
MOTA MOTA	900 901	CO	ARG ARG	96	13.114	60.072	109.135	1.00	13.09	0
MOTA	902	N	GLY	97	11.668	58.411 57.480	109.592 108.716	1.00	19.94 20.45	0 0
MOTA	904 905	CA C	GLY GLY	97 97	12.359 12.412	57.920	107.269	1.00	23.76	, 0
MOTA MOTA	906	0	GLY	97	11.516	58.617	106.773		84.64	0
MOTA	907	N	LYS	98	13.460 13.698	57.469 57.806	106.584 105.182		47.57 46.79	Ö
M OTA MOTA	909 910	CA CB	LYS LYS	98 98	15.147	57.462	104.832		31.45	0
UI OU	710	CD	5.5							

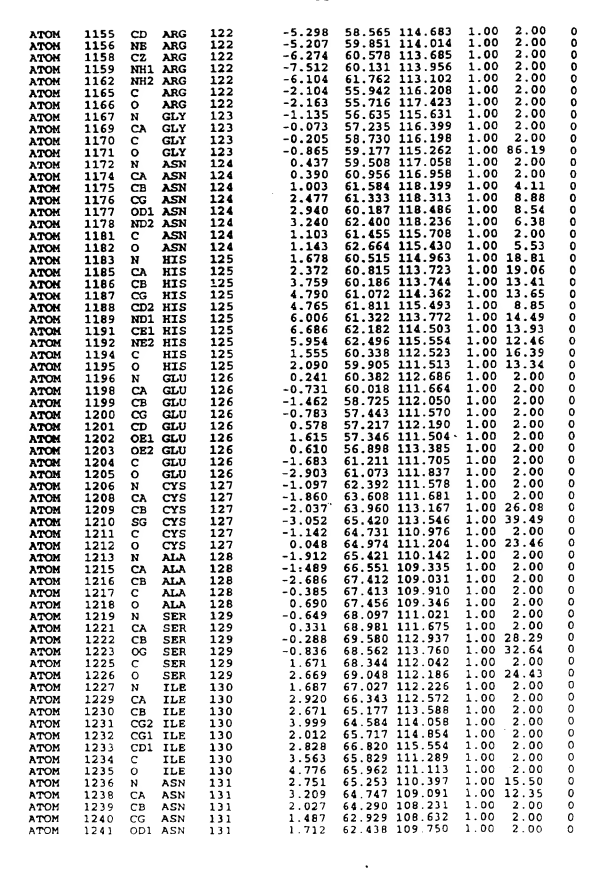
ATOM	911	CG	LYS	98	16.169	58.397 105.487	1.00 30.73	0
ATOM	912	CD	LYS	98	17.606	58.061 105.087	1.00 32.79	0
MOTA	913	CE	LYS	98	18.605 20.036	59.089 105.627 58.743 105.315	1.00 30.08 1.00 31.54	0
MOTA	914 918	NZ C	LYS LYS	98 98	12.741	57.196 104.146	1.00 44.78	Ö
MOTA MOTA	919	0	LYS	98	12.613	57.707 103.040	1.00 31.43	0
ATOM	920	N	GLN	99	12.059	56.120 104.517	1.00 2.00	0
MOTA	922	CA	GLN	99	11.132 11.654	55.430 103.639 54.023 103.337	1.00 2.00 1.00 11.21	0
MOTA	923	CB	GLN GLN	99 99	12.945	53,993 102.552	1.00 6.66	ŏ
MOTA MOTA	924 925	CG CD	GLIN	99	13.361	52.586 102.177	1.00 11.01	Ö
MOTA	926	OE1	GLN	99	13.802	51.805 103.031	1.00 10.44	0
MOTA	927	NE2	GLN	99	13.229	52.247 100.895 55.328 104.245	1.00 8.18 1.00 2.00	0
ATOM	930	C	GLN	99 99	9.741 9.177	54.238 104.317	1.00 6.66	Ö
MOTA	931 932	O N	GLN SER	100	9.178	56.460 104.657	1.00 24.01	ŏ
MOTA MOTA	934	CA	SER	100	7.839	56.475 105.257	1.00 24.01	0
MOTA	935	CB	SER	100	7.481	57.897 105.739	1.00 2.00	0
ATOM	936	OG	SER	100	8.479 6.749	58.490 106.556 55.987 104.272	1.00 2.00 1.00 24.01	0
MOTA	938	Č	SER	100 100	5.703	55.463 104.680	1.00 2.00	ŏ
ATOM ATOM	939 940	O N	SER Leu	101	7.015	56.153 102.977	1.00 27.83	0
ATOM	942	CA	LEU	101	6.084	55.786 101.913	1.00 27.83	0
ATOM	943	CB	LEU	101	6.551	56.416 100.599 57.224 99.721	1.00 3.66 1.00 3.66	0
ATOM	944	CG	LEU	101	5.593 4.209	57.224 99.721 56.591 99.749	1.00 3.66	ŏ
MOTA	945 946	CD1	LEU LEU	101 101	5.532	58.653 100.208	1.00 3.66	Ō
MOTA MOTA	947	CDZ	LEU	101	5.852	54.284 101.689	1.00 27.83	0
ATOM	948	Ö	LEU	101	4.731	53.851 101.467	1.00 3.66	0
MOTA	949	N	GLU	102	6. 901 6. 71 3	53.480 101.725 52.055 101.490	1.00 2.00 1.00 2.00	ŏ
ATOM	951	CA	GLU GLU	102 102	7.976	51.470 100.870	1.00 13.98	ŏ
MOTA MOTA	952 953	CB CG	GLU	102	9.211	52.208 101.294	1.00 13.98	0
MOTA	954	œ	GLU	102	10.116	52.539 100.136	1.00 13.98	0
ATOM	955	OE1		102	10.181	53.727 99.737 51.599 99.643	1.00 13.98 1.00 13.98	0
ATOM	956	OE2		102 102	10.772 6.307	51.324 102.763	1.00 2.00	ŏ
MOTA MOTA	957 958	CO	GLU	102	5.686	50,263 102.729	1.00 13.98	0
MOTA	959	N	THR	103	6.664	51.901 103.897	1.00 2.00 1.00 2.00	0
ATOM	961	CA	THR	103	6.293	51.330 105.173 52.098 106.309	1.00 2.00 1.00 2.00	Ö
MOTA	962	CB	THR	103 103	6.923 8.313	52.297 106.028	1.00 2.00	Ö
MOTA MOTA	963 965	OG1 CG2		103	6.758	51.329 107.599	1.00 2.00	0
ATOM	966	C	THR	103	4.775	51.440 105.296	1.00 2.00	0
ATOM	967	0	THR	103	4.081	50.428 105.394 52.667 105.257	1.00 2.00 1.00 2.00	0
MOTA	968	N	ILE	104	4.256 2.824	52.667 105.257 52.873 105.363	1.00 2.00	ŏ
MOTA	970 971	CA CB	ILE	104 104	2.486	54.389 105.300	1.00 12.72	0
MOTA MOTA	972	CG2		104	2.856	54.965 103.956	1.00 12.35	0
MOTA	973		ILE	104	0.996	54.618 105.548	1.00 13.84 1.00 12.35	0
MOTA	974	CD1		104	0.459 2.074	53.940 106.787 52.062 104.294	1.00 2.00	ŏ
MOTA	975 976	C O	ILE	104 104	0.975	51.581 104.544	1.00 18.28	0
MOTA MOTA	977	N	CYS	105	2.690	51.868 103.127	1.00 23.45	0
ATOM	979	CA	CYS	105	2.080	51.092 102.042	1.00 21.74 1.00 20.76	0
MOTA	980	CB	CYS	105	2.722 2.010	51.458 100.708 52.958 99.957	1.00 18.03	ō
ATOM	981	SG	CYS	105 105	2.061	49.563 102.214	1.00 17.08	0
MOTA MOTA	982 983	C O	CYS	105	1.114	48.915 101.781	1.00 20.76	0
ATOM	984	N	LEU	106	3.089	48.982 102.833	1.00 2.00 1.00 2.00	0
MOTA	986	CA	LEU	106	3.124	47.526 103.073 47.042 103.495	1.00 2.00	Ö
MOTA	987	CB	LEU	106 106	4.519 4.680	45.544 103.802	1.00 2.00	0
MOTA MOTA	988 989	CG	LEU LEU	106	4.200	44.694 102.639	1.00 2.00	0
MOTA	990		LEU	106	6.133	45.247 104.053	1.00 2.00	0
ATOM			LEU	106	2.154	47.179 104.178	1.00 2.00	
	991	С				46 000 104 192	1.00 2 00	[]
MOTA MOTA	991 992 993	0 0	LEU	106 107	1.589 1.992	46.088 104.192 48.107 105.116	1.00 2.00 1.00 2.00	0

ATOM AOTA MOTA MOTA MOTA MOTA MOTA MOTA	995 996 997 998 999 1000 1001 1002 1004 1005 1006 1007 1008	CA LEU CB LEU CG LEU CD1 LEU CD2 LEU C LEU O LEU N LEU CA LEU CB LEU CG LEU CG LEU CD1 LEU CD1 LEU	107 107 107 107 107 107 108 108 108 108	1.078 1.347 2.761 3.106 2.868 -0.339 -1.153 -0.623 -1.953 -2.020 -2.103 -2.244 -3.281	47.922 106.226 48.967 107.319 48.857 107.911 50.016 108.815 47.535 108.638 48.035 105.683 47.125 105.873 49.126 104.971 49.340 104.394 50.666 103.637 51.925 104.499 53.164 103.618 51.792 105.433	1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1009 1010 1011 1013 1014 1015 1016 1017 1020 1021 1022 1023 1024	C LEU O LEU N ALA CA ALA CB ALA C ALA O ALA N TYR CA TYR CB TYR CG TYR CD1 TYR CC1 TYR CC1 TYR CC1 TYR	108 108 109 109 109 110 110 110 110	-2.352 -3.533 -1.366 -1.608 -0.336 -2.091 -3.146 -1.311 -1.625 -0.495 0.674 1.966 3.053	48.206 103.455 47.846 103.369 47.643 102.762 46.539 101.839 46.209 101.087 45.316 102.605 44.746 102.305 44.929 103.609 43.781 104.452 43.564 105.438 42.771 104.922 43.235 105.092 42.474 104.706 41.517 104.340	1.00 2.00 1.00 2.00 1.00 2.00 1.00 33.06 1.00 2.00 1.00 24.92 1.00 5.11 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1025 1026 1027 1029 1030 1031 1033 1034 1035 1036 1037 1038	CE2 TYR CZ TYR OH TYR C TYR O TYR N LYS CA LYS CB LYS CG LYS CD LYS CD LYS CE LYS	110 110 110 110 111 111 111 111 111 111	1.578 2.859 3.963 -2.948 -3.663 -3.265 -4.508 -4.619 -5.942 -6.085 -7.410 -7.643 -5.698	40.743 103.950 41.230 104.139 40.483 103.779 43.940 105.215 42.961 105.441 45.168 105.618 45.430 106.333 46.904 106.708 47.262 107.393 46.501 108.685 46.743 109.354 45.735 110.428 45.051 105.465	1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1043 1044 1046 1047 1048 1059 1051 1052 1053 1055 1056 1057	O LYS N ILE CA ILE CB ILE CG2 ILE CG1 ILE CD1 ILE O ILE N LYS CA LYS CB LYS CG LYS	111 112 112 112 112 112 112 113 113 113	-6.655° -5.624 -6.651 -6.361 -7.414 -6.339 -5.857 -6.706 -7.783 -5.537 -5.537 -4.001 -3.852	44.431 105.951 45.455 104.192 45.192 103.177 45.978 101.875 45.656 100.814 47.482 102.166 48.336 101.012 43.706 102.829 43.156 102.568 43.067 102.816 41.649 102.513 41.285 102.181 39.909 101.596	1.00 2.00 1.00 2.00 1.00 25.04 1.00 25.04 1.00 25.04 1.00 25.04 1.00 25.04 1.00 25.04 1.00 2.00 1.00 2.00 1.00 8.72 1.00 8.72 1.00 8.72	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1058 1059 1060 1064 1065 1066 1069 1070 1071 1072 1073 1074	CD LYS CE LYS NZ LYS O LYS O LYS N TYR CA TYR CB TYR CG TYR CD1 TYR CD1 TYR CD1 TYR CD2 TYR CC2 TYR CC4 TYR CC7 TYR	113 113 113 113 114 114 114 114 114 114	-2.780 -2.618 -3.952 -5.987 -6.620 -5.744 -6.226 -5.122 -4.138 -3.027 -2.111 -4.307 -3.395 -2.296 -1.378	39.871 100.521 38.467 99.975 37.873 99.637 40.780 103.667 39.742 103.435 41.200 104.906 40.455 106.068 39.580 106.666 38.988 105.689 39.715 105.278 39.191 104.397 37.709 105.188 37.168 104.302 37.917 103.908 37.404 103.013	1.00 8.72 1.00 8.72 1.00 8.72 1.00 2.00 1.00 8.72 1.00 9.02 1.00 12.47 1.00 12.47 1.00 12.47 1.00 12.47 1.00 12.47 1.00 12.47 1.00 12.47 1.00 12.47 1.00 12.47	

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ATON ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1078 1079 1080 1081 1082 1083 1084 1085 1089 1099 1099 1099 1109 1109 1109 1109	ND2 C O N CA CB CG CD1 CD2 CE1 CE2 CZ C	TYR PRO PRO PRO PRO PRO GLU GLU GLU GLU ASN ASN ASN PHE	114 115 115 115 115 115 116 116 116 116 116	-6.729 -6.729 -7.928 -8.843 -8.8488 -9.640 -8.5017 -9.275 -9.477 -10.577 -11.6739 -13.0274 -5.820 -5.457 -6.328 -4.613 -3.135 -1.925 -1.995 -1.995 -1.995 -1.995 -1.995 -1.995	41.417 107 41.638 108 41.985 106 41.845 105 42.912 107 43.227 107 43.117 105 42.837 110 41.181 109 40.538 110 39.469 110 39.726 109 40.728 100 41.641 109 40.599 111 39.902 111 39.902 111 39.902 111 39.902 111 39.952 111 38.220 110 37.351 109 36.071 111 40.372 111 39.952 111 40.372 111 39.952 111 40.372 111	.169 .968 .824 .3215 .235 .235 .235 .235 .235 .235 .235 .23	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	34.30 24.86 34.30 24.86 24.86 34.30 24.86 2.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM	1119 1120 1122	O N CA	PHE PHE PHE	118 119 119	-3.874 -4.467		.240 .721		12.11	0 0 0
ATOM ATOM	1123 1124	CB CG	PHE PHE	119 119	-5.174 -6.229	44.355 113	.889	1.00	2.00	0
MOTA	1125 1126	CD1	PHE	119 119	-5.903 -7.556	43.008 113 44.716 113	.975 .655	1.00 1.00	2.00 2.00	0
atom Mota	1127	CE1	PHE	119	-6.881	42.035 113	.829	1.00 1.00	2.00 2.00	0
MOTA MOTA	1128 1129	CE2	PHE PHE	119 119	-8.546 -8.209	42.410 113	.506 .592	1.00	2.00	0
MOTA	1130	С	PHE	119	-3.472	46.806 112	.905 .346	1.00	12.11 2.00	0
MOTA MOTA	1131 1132	N O	PHE LEU	119 120	-2.342 -3.893	48.008 112	.537	1.00	2.00	0
ATOM	1134	CA	LEU	120	-3.043	49.179 112 49.809 111	.672	1.00 1.00	2.00 2.00	0
MOTA	1135 1136	CB CG	LEU	120 120	-2.770 -2.127	48.910 110	.259	1.00	2.00	0
MOTA MOTA	1137	CD1	LEU	120	-2.147	49.623 108	.924	1.00	2.00	0
MOTA	1138	CD2	LEU	120	-0.731 -3.766	48.539 110 50.176 113	559	1. 0 0 1. 0 0	2.00	ŏ
MOTA MOTA	1139 1140	С О	LE U	120 120	-4.963	50,401 113	. 37 5	1.00	2.00	0
MOTA	1141	N	LEU	121	-3.069		.542	1.00 1.00	2. 0 0 2. 0 0	0
MOTA	1143	CA	LEU	121	-3.670 -3.351		.415 .890	1.00	2.00	ŏ
MOTA MOTA	1144 1145	CB CG	LEU	121 121	-4.142	50.320 117	.598	1.00	2.00	0
MOTA	1146	CD1	LBU	121	-3.648		.012	1.00 1.00	2.00	0 0
ATOM	1147	CD2	LEU	121 121	-5.609 -3.106	50.657 117 53.060 115		1.00	2.00	0
MOTA MOTA	1148 1149	Ö	LEU	121	-2.213	53.148 114	.166	1.00	2.00	0
MOTA	1150	N	ARG	122	-3.631	54.118 115 55.434 115	.592 . 251	1.00	2.00	0
MOTA	1152 1153	CA CB	ARG ARG	122 122	-3.162 -4.336	56.404 115	.224	1.00	2.00	ő
MOTA MOTA	1153	CC	ARG	122	-4.047	57.765 114	.619	1.00	2.00	0

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				0.245	62.313 107.718	1.00 2.00	0
MOTA		ND2 ASN	131	0.745	65.914 108.374	1.00 28.24	ŏ
MOTA	1245	C ASN	131	3.849	65.854 107.951	1.00 2.00	ŏ
MOTA	1246	o asn	131	5.001	66.971 108.255	1.00 2.00	ŏ
MOTA	1247	n arg	132	3.051	68.234 107.627	1.00 2.00	ŏ
MOTA	1249	CA ARG	132	3.386	69.269 108.006	1.00 22.25	ŏ
MOTA	1250	CB ARG	132	2.327	70.673 107.477	1.00 26.48	Ö
MOTA	1251	CG ARG	132	2.505	70.734 105.962	1.00 27.28	ŏ
MOTA	1252	CD ARG	132	2.505	70.533 105.434	1.00 36.02	ŏ
MOTA	1253	ne arg	132	3.849	71.479 104.852	1.00 34.03	ŏ
ATOM	125 5	CZ ARG	132	4.577	72.697 104.711	1.00 30.29	ŏ
MOTA	1256	NH1 ARG	132	4.080	71.211 104.424	1.00 37.77	Ö
MOTA	1259	NH2 ARG	132	5.808	68.731 107.990	1.00 2.00	ŏ
MOTA	1262	C ARG	132	4.768	69.477 107.225	1.00 27.60	ŏ
MOTA	1263	O ARG	132	5.359	68.324 109.127	1.00 2.00	Ö
MOTA	1264	N ILE	133	5.314	68.801 109.457	1.00 2.00	ŏ
MOTA	1266	CA ILE	133	6.652	69.644 110.746	1.00 20.12	ŏ
MOTA	1267	CB ILE	133	6.652	71.055 110.441	1.00 19.34	Õ
MOTA	1268	CG2 ILE	133	6.215	68.999 111.800	1.00 21.76	Ö
MOTA	1269	CG1 ILE	133	5.761	69.792 113.065	1.00 22.74	ŏ
MOTA	1270	CD1 ILE	133	5.687	67.779 109.553	1.00 2.00	ō
MOTA	1271	C ILE	133	7.794	68.073 109.127	1.00 21.74	ŏ
MOTA	1272	O ILE	133	8.915	66.583 110.080	1.00 2.00	ŏ
MOTA	1273	n tyr	134	7.521	65.567 110.270	1.00 2.00	ŏ
MOTA	1275	CA TYR	134	8.571 8.330	64.766 111.561	1.00 27.01	0
MOTA	1276	CB TYR	134	8.270	65.648 112.767	1.00 19.94	0
MOTA	1277	CG TYR	134 134	7.092	65.789 113.486	1.00 23.86	0
ATOM	1278	CD1 TYR	134	7.016	66.651 114.557	1.00 24.86	0
ATOM	1279 1280	CE1 TYR	134	9.380	66.392 113.160	1.00 24.55	0
MOTA	1281	CE2 TYR	134	9.312	67.254 114.228	1.00 23.33	0
MOTA MOTA	1282	CZ TYR	134	8.128	67.384 114.925	1.00 28.93	0
ATOM	1283	OH TYR	134	8.043	68.259 115.984	1.00 27.73	0
ATOM	1285	C TYR	134	8.823	64.606 109.139	1.00 2.00	ŏ
MOTA	1286	O TYR	134	8.986	63.404 109.368	1.00 24.62	Ö
ATOM	1287	N GLY	135	8.847	65.127 107.922	1.00 11.68 1.00 10.40	ŏ
ATOM	1289	CA GLY	135	9.134	64.275 106.787	1.00 9.03	ŏ
MOTA	1290	C GTX	135	8.028	63.772 105.882 64.127 104.713	1.00 20.70	ŏ
MOTA	1291	O GLY	135	B.010	64.127 104.713 62.963 106.393	1.00 2.00	Õ
MOTA	1292	N PHE	136	7.110 6.0 6 2	62.412 105.549	1.00 2.00	0
MOTA	1294	CA PHE	136	4.951	61.757 106.347	1.00 2.00	0
MOTA	1295	CB PHE	136	4.091	60.831 105.524	1.00 2.00	0
MOTA	1296	CG PHE	136	4.664	59.734 104.870	1.00 2.00	0
MOTA	1297	CD1 PHE	136 136	2.713	61.036 105.424	1.00 2.00	0
ATOM	1298	CD2 PHE	136	3.882	58.846 104.131	1.00 2.00	0
ATOM	1299	CE1 PHE CE2 PHE	136	1.915	60.155 104.688	1.00 2.00	0
ATOM	1300	_	136	2.507	59.050 104.037	1.00 2.00	0
MOTA	1301 1302	CZ PHE C PHE	136	5.421	63.358 104.563	1.00 2.00	0
MOTA MOTA	1302	O PHE	136	5.121	62.926 103.445	1.00 2.00	0
ATOM	1304	N TYR	137	5. 19 9	64.616 104.940	1.00 2.00 1.00 2.00	Ö
MOTA	1306	CA TYR	137	4.606	65.530 103.977		ŏ
MOTA	1307	CB TYR	137	4.243	66.859 104.598		ŏ
MOTA	1308	CG TYR	137	3.886	67.921 103.564	1.00 2.00 1.00 2.00	Ö
MOTA	1309	CD1 TYR	137	2.550	68.203 103.257	1.00 2.00	ō
MOTA	1310	CE1 TYR	137	2.228	69.196 102.327 68.666 102.901	1.00 2.00	Ö
ATOM	1311	CD2 TYR	137	4.893		1.00 2.00	0
MOTA	1312	CE2 TYR	137	4.576	69.644 101.985 69.903 101.707	1.00 2.00	0
ATOM	1313	CZ TYR	137	3.247	70.893 100.823	1.00 2.00	0
MOTA	1314	OH TYR	137	2.920 5.566	65.785 102.822	1.00 2.00	0
MOTA	1316	C TYR	137	5.187	65.669 101.646	1.00 2.00	O
MOTA	1317	O TYR	137	6.802	66.153 103.155	1.00 2.00	0
ATOM	1318	N ASP	138 138	7.822	66.439 102.147	1.00 2.00	0
MOTA	1320	CA ASP CB ASP	138	9.102	66.971 102.810	1.00 28.59	0
MOTA	1321		138	8.832	6B.173 103.705	1.00 31.85	0
MOTA	1322 1323	CG ASP	138	8.999	69.327 103.255	1.00 34.04	0
MOTA MOTA	1323	OD2 ASP	138	8.432	67.963 104.867	1.00 31.26	0
MOTA	1325	C ASP	138	8.118	65.203 101.308	1.00 2.00	0

								4 00 07 30	_
MOTA	1326	0	ASP	138	8.322	65.309	100.102	1.00 27.39	0
MOTA	1327	N	GLU	139	8.097	64.030	101.926	1.00 31.40	0
MOTA	1329	CA	GLU	139	8.366	62.797	101.200	1.00 28.32	0
		_		139	8.380		102.149	1.00 2.00	0
MOTA	1330	CB	GLU		8.965		101.526	1.00 4.78	Õ
MOTA	1331	CG	GLU	139		50.301	102.525	1.00 3.01	ŏ
MOTA	1332	CD	GLU	139	9.167				
MOTA	1333	OE1	GLU	139	9.135		102.087	1.00 2.00	0
MOTA	1334	OE2	GLU	139	9.372	59.518	103.738	1.00 4.61	0
MOTA	1335	C	GLU	139	7.323	62.572	100.117	1.00 30.47	0
			GLU	139	7.660	62.419	98.941	1.00 2.00	0
MOTA	1336	0			6.055		100.511	1.00 12.32	Ō
ATOM	1337	N	CYS	140			99.553	1.00 3.99	ŏ
MOTA	1339	CA	CYS	140	4.981	62.355			
MOTA	1340	CB	CYS	140	3.625		100.221	1.00 18.56	0
ATOM	1341	SG	CYS	140	3.313	61.308	101.477	1.00 23.54	0
	1342	C	CYS	140	5.128	63.346	98.416	1.00 12.32	0
ATOM			CYS	140	5.117	62.961	97.257	1.00 21.26	0
MOTA	1343	0				64.616	98.763	1.00 41.55	ō
MOTA	1344	N	LYS	141	5.305				
MOTA	1346	CA	LYS	141	5.446	65.696	97.786	1.00 40.86	0
MOTA	1347	CB	LYS	141	5.655	67.023	98.531	1.00 38.66	0
MOTA	1348	CG	LYS	141	5.853	68.246	97.646	1.00 32.19	0
	1349	CD	LYS	141	5.886	69.522	98.468	1.00 35.28	0
ATOM				141	5.895	70.750	97.584	1.00 36.09	0
MOTA	1350	CE	LYS					1.00 40.44	ŏ
MOTA	1351	NZ	LYS	141	5.552	71.948	98.380		
ATOM	1355	С	LYS	141	6.586	65.467	96.791	1.00 40.27	0
ATOM	1356	0	LYS	141	6.431	65.642	95.582	1.00 28.60	0
ATOM	1357	N	ARG	142	7.731	65.073	97.319	1.00 23.84	0
				142	8.912	64.834	96.512	1.00 23.84	0
ATOM	1359	CA	ARG		10.097	64.566	97.444	1.00 22.75	Ŏ
MOTA	1360	CB	ARG	142				1.00 24.10	ŏ
ATOM	1361	CG	ARG	142	11.368	64.101	96.778		
ATOM	1362	CD	ARG	142	12.474	64.073	97.799	1.00 33.58	0
MOTA	1363	NE	ARG	142	12.099	63.283	98.963	1.00 37.62	0
	1365	CZ	ARG	142	12.464	62.018	99.150	1.00 46.30	0
MOTA					13.224	61.404	98.249	1.00 44.23	0
MOTA	1366	NH1		142			100.234	1.00 42.22	ŏ
MOTA	1369	NH2	ARG	142	12.060	61.363			ŏ
MOTA	1372	С	ARG	142	8.732	63.689	95.519`	1.00 23.84	U
		•	M						
					8.995	63.838	94.330	1.00 24.93	0
ATOM	1373	0	ARG	142	8.995	63.838			
ATOM ATOM	1373 1374	O N	ARG ARG	142 143	8.995 8.268	63.838 62.550	94.330 95.999	1.00 24.93 1.00 2.00	0
ATOM ATOM ATOM	1373 1374 1376	CA N	ARG ARG ARG	142 143 143	8.995 8.268 8.104	63.838 62.550 61.413	94.330 95.999 95.125	1.00 24.93 1.00 2.00 1.00 2.00	0 0 0
MOTA MOTA MOTA	1373 1374 1376 1377	O N CA CB	ARG ARG ARG ARG	142 143 143 143	8.995 8.268 8.104 8.267	63.838 62.550 61.413 60.134	94.330 95.999 95.125 95.941	1.00 24.93 1.00 2.00 1.00 2.00 1.00 2.86	0 0 0
ATOM ATOM ATOM	1373 1374 1376 1377 1378	O N CA CB CG	ARG ARG ARG ARG	142 143 143 143 143	8.995 8.268 8.104 8.267 9.686	63.838 62.550 61.413 60.134 59.941	94.330 95.999 95.125 95.941 96.455	1.00 24.93 1.00 2.00 1.00 2.00 1.00 2.86 1.00 2.86	0 0 0 0
MOTA MOTA MOTA	1373 1374 1376 1377	O N CA CB	ARG ARG ARG ARG	142 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792	63.838 62.550 61.413 60.134 59.941 58.874	94.330 95.999 95.125 95.941 96.455 97.530	1.00 24.93 1.00 2.00 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64	0 0 0 0
ATOM MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379	O N CA CB CG CD	ARG ARG ARG ARG	142 143 143 143 143	8.995 8.268 8.104 8.267 9.686	63.838 62.550 61.413 60.134 59.941	94.330 95.999 95.125 95.941 96.455 97.530 97.049	1.00 24.93 1.00 2.00 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64 1.00 4.03	0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380	O N CA CB CG CD NE	ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792	63.838 62.550 61.413 60.134 59.941 58.874	94.330 95.999 95.125 95.941 96.455 97.530	1.00 24.93 1.00 2.00 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91	0000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379 1380 1382	O N CA CB CG CD NE CZ	ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439	94.330 95.999 95.125 95.941 96.455 97.530 97.049	1.00 24.93 1.00 2.00 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64 1.00 4.03	0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380 1382 1383	O N CA CB CG CD NE CZ NH1	ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13	0000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386	O N CA CB CG CD NE CZ NH1 NH2	ARG ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97	0000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389	O N CA CB CG CD NE CZ NH1	ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796	63.838 62.550 61.413 60.134 59.941 57.527 56.527 56.527 55.257 61.411	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	00000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386	O N CA CB CG CD NE CZ NH1 NH2	ARG ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.203 6.796 6.707	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.439 57.527 61.411 60.818	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 93.295	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 12.04	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389	O N CA CB CG CD NE CZ NH1 NH2 C	ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 55.257 61.411 60.818 62.097	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 93.295 94.897	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 12.04 1.00 2.00	0 0 0 0 0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389 1390 1391	O N CA CB CG CD NE CZ NH1 NH2 C O N	ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	142 143 143 143 143 143 143 143 143 143	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.203 6.796 6.707	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.439 57.527 61.411 60.818	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 93.295	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 12.04 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1399 13991 1393	O N CA CB CG CD NE CZ NH1 NH2 C O N CA	ARG ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR	142 143 143 143 143 143 143 143 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.097 62.120	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 93.295 94.897	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 12.04 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1390 1393 1393	O N CA CB CCZ NH1 NH2 C O N CA CB	ARG ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR	142 143 143 143 143 143 143 143 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 60.818 60.8150 61.150	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.897 94.897 94.274	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 12.04 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1390 1391 1393 1394 1395	O N CA CB CC NH1 NH2 C O N CA CB CC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR	142 143 143 143 143 143 143 143 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.689	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.897 94.274 95.021 94.953	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389 1390 1391 1393 1395 1396	O N CA CB CCD NH1 NH2 C O N CA CB CCD 1	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR	142 143 143 143 143 143 143 143 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946	63.838 62.550 61.413 60.134 59.941 58.527 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.689 59.203	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 94.295 94.295 94.274 95.021 95.716	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389 1390 1391 1393 1395 1396 1397	O N CA CB CCD NE CZ NH1 NH2 C O N CA CCB CCD1 CE1	ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR	142 143 143 143 143 143 143 143 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.509 3.509	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.097 62.120 619.689 59.203 57.885	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 94.295 94.295 94.274 95.021 95.635	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389 1390 1391 1393 1395 1396	O N CA CB CCD NH1 NH2 C O N CA CB CCD 1	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR	142 143 143 143 143 143 143 143 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.509 3.509	63.838 62.550 61.413 60.134 59.941 57.527 56.527 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.203 57.885 58.804	94.330 95.999 95.125 95.941 96.455 97.530 97.049 99.115 97.287 94.367 94.274 95.021 94.975 94.975 94.975 94.975	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.66 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1389 1391 1393 1394 1395 1397 1398	O N CA CB CCD NH1 NH2 C O N CA CB CCD 1 CD1 CD2	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR	142 143 143 143 143 143 143 143 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.509 3.509	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.097 62.120 619.689 59.203 57.885	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 94.295 94.295 94.274 95.021 95.635	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1380 1382 1383 1386 1399 1391 1393 1394 1395 1397 1398 1399	O N CA CB CCD NH1 NH2 C O N CA CB CCD CD1 CD2 CD2 CD2	ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 143 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633	63.838 62.550 61.413 60.134 59.941 57.527 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.689 59.203 57.885 58.804 57.483	94.330 95.999 95.125 95.941 96.455 97.530 97.049 97.816 99.115 97.287 94.367 94.274 95.021 94.953 95.635 94.102 94.017	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.66 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1399 1395 1396 1397 1398 1399 1400	O N CA CCC CCC NH1 NH2 C O N CA CCC CCC CCC CCC CCC CCC CCC CCC CC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633 4.683	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.819 62.120 61.150 59.689 59.203 57.885 58.804 57.034	94.330 95.999 95.125 95.945 96.455 97.049 97.816 99.115 97.287 94.367 94.274 95.021 94.953 95.716 95.632 94.017 94.786	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1399 1399 1395 1397 1398 1399 1400 1401	O N CA CB CCD NH1 NH2 C O N CA CB CCD1 CCD2 CCD CCD2 CCD2 CCC CCC CCC CCC CCC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 143 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.707 5.707 5.707 4.459 3.509 3.902 4.946 5.336 3.248 3.4683 5.113	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.818 60.818 60.818 59.203 57.885 58.804 57.034 55.738	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.274 95.021 94.953 95.635 94.102 94.703	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1399 1399 1395 1397 1398 1398 1399 1400 1401 1403	O N CA CB CCD NH1 NH2 C O N CA CB CCD CE1 CCC CCC CCC CCC CCC CCC CCC CCC CCC	ARG ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.683 5.113 3.789	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.0120 61.150 59.689 59.203 57.885 58.804 57.483 57.034 55.738 63.502	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.897 94.274 95.021 94.953 95.635 94.102 94.703 94.703 94.703	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1399 1399 1395 1396 1397 1398 1399 1400 1403 1404	O N CA CB CCD NH1 NH2 C O N CA CB CCD1 CCD2 CCD CCD2 CCD2 CCC CCC CCC CCC CCC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633 4.683 5.113 3.789 4.129	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.689 59.203 57.885 58.804 57.483 57.034 63.502 64.316	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.295 94.897 94.295 94.953 95.635 94.7194.7883 94.769 93.297	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1380 1382 1383 1386 1389 1399 1399 1395 1397 1398 1398 1399 1400 1401 1403	O N CA CB CCD NH1 NH2 C O N CA CB CCD CE1 CCC CCC CCC CCC CCC CCC CCC CCC CCC	ARG ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.902 4.946 5.336 3.248 3.633 4.633 4.129 2.827	63.838 62.550 61.413 60.134 59.941 58.527 56.527 56.527 61.411 60.818 62.097 62.120 61.150 59.203 57.885 58.804 57.483 57.738 63.750	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.295 94.295 94.295 94.271 94.716 94.716 94.716 94.716 94.716 94.716 94.716 95.916	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1380 1382 1388 1389 1399 1399 1399 1399 1400 1403 1404 1405	O N CA CB CCD NE CZ NH1 NH2 C O N C CB CCD C C C C C C C C C C C C C C C	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633 4.683 5.113 3.789 4.129	63.838 62.550 61.413 60.134 59.941 58.527 56.527 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.203 57.885 58.804 57.483 57.034 55.738 63.750 63.750 65.000	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.295 94.897 94.295 94.953 95.635 94.7194.7883 94.769 93.297	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1382 1383 1386 1389 1399 1399 1400 1401 1403 1404 1405 1407	O N CAB CCD NE C NH1 NH2 C O N CAB CCD C C C C C C C C C C C C C C C C C	ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	142 143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.795 4.459 3.509 3.902 4.946 5.336 3.633 4.683 5.113 3.729 2.827 2.081	63.838 62.550 61.413 60.134 59.941 58.527 56.527 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.203 57.885 58.804 57.483 57.034 55.738 63.750 63.750 65.000	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.295 94.295 94.295 94.271 94.716 94.716 94.716 94.716 94.716 94.716 94.716 95.058	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1378 1379 1382 1383 1386 1389 1399 1399 1399 1400 1401 1403 1404 1405 1407 1408	O N CA CG CC	ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.795 3.509 3.509 3.509 3.548 3.633 4.683 5.113 3.789 4.129 2.827 2.081 1.159	63.838 62.550 61.413 60.134 59.941 58.527 56.527 56.527 55.257 61.411 60.818 62.097 62.120 61.150 59.203 57.885 58.804 57.483 57.034 55.738 63.750 63.750 65.000 65.078	94.330 95.999 95.125 95.941 96.455 97.530 97.816 97.287 94.274 95.025 94.271 94.7023 95.632 94.703 94.703 94.703 94.703 94.703 94.703 94.703 94.703 94.703 94.703 95.0867	1.00 24.93 1.00 2.00 1.00 2.86 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1380 1388 1389 1399 1399 1399 1399 1400 1403 1404 1405 1408 1409	O N CAB CCD NH2 C O N CAB CCD11 CCE2 CH C O N CAB CCD1 CCE2 CH C O N CAB CCC CCC CCC CCC CCC CCC CCC CCC CCC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	14333333333333444444444444444444444444	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.782 9.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633 4.683 5.113 3.789 4.129 2.8081 1.159 0.255	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.897 62.120 61.150 59.689 59.203 57.885 57.885 57.034 57.738 63.502 64.316 63.750 65.078 63.861	94.330 95.999 95.125 95.945 96.455 97.049 97.816 99.115 97.287 94.276 94.271 94.271 94.271 94.271 94.716 95.632 94.1017 94.7169 94.7169 94.7169 94.7169 94.7169 94.7169 95.0867 95.0867 97.93.748	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1380 1382 1383 1386 1389 1399 1399 1399 1400 1403 1404 1405 1407 1409 1410	ON CAB CCD NH2 CON CAB CCD1 CCC CC ON CAB CCD1 CCC CCC CCC CCC CCC CCC CCC CCC CC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR ASN ASN ASN ASN	143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.707 5.707 5.707 4.459 3.509 3.902 4.946 5.336 3.248 3.683 4.683 5.113 3.789 4.129 2.827 2.081 1.159 0.255 -0.823	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.818 60.818 62.120 61.150 59.689 59.203 57.885 58.804 57.034 55.738 63.502 64.316 63.750 65.000 65.000 65.000 65.000 65.8861 63.826	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.274 95.053 94.716 95.632 94.716	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1380 1388 1389 1399 1399 1399 1399 1400 1403 1404 1405 1408 1409	O N CAB CCD NH2 C O N CAB CCD11 CCE2 CH C O N CAB CCD1 CCE2 CC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.707 5.309 3.509 3.902 4.946 5.336 3.248 3.683 5.113 3.789 4.129 2.827 2.081 1.159 0.255 -0.823 0.702	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.120 61.150 59.689 59.203 57.885 58.804 57.438 63.750 64.316 63.750 65.078 65.078 65.078 65.078 65.078 65.078 65.078 65.078 65.078 65.078	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.274 95.053 94.021 94.763 94.7169 94.769 94.769 94.769 94.769 94.769 94.769 94.769 94.769 95.0869 97.869	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1380 1382 1383 1386 1389 1399 1399 1399 1400 1403 1404 1405 1407 1409 1410	ON CAB CCD NH2 CON CAB CCD1 CCC CC ON CAB CCD1 CCC CCC CCC CCC CCC CCC CCC CCC CC	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR ASN ASN ASN ASN	143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633 4.683 5.113 3.789 4.129 2.827 2.081 1.159 0.702 1.229	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.818 62.097 61.150 59.689 59.203 57.885 58.804 57.483 57.034 57.034 63.750 64.316 63.750 65.000 65.078 63.861 63.851 63.851 63.851 63.851 63.851	94.330 95.999 95.125 95.941 96.4550 97.049 97.816 99.115 97.287 94.3297 94.271 94.953 94.023 94.023 94.071 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1382 1388 1389 1399 1399 1399 1399 1399 1403 1404 1405 1407 1408 1409 1411 1414	O N CA CCC CCC NH1 NH2 C O N CA CCC CCC CCC CCC CCC CCC CCC CCC CC	ARG ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	14333333333344444444444444444444444444	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.707 5.309 3.509 3.902 4.946 5.336 3.248 3.683 5.113 3.789 4.129 2.827 2.081 1.159 0.255 -0.823 0.702	63.838 62.550 61.413 60.134 59.941 58.874 57.527 56.439 56.527 55.257 61.411 60.818 62.120 61.150 59.689 59.203 57.885 58.804 57.438 63.750 64.316 63.750 65.078 65.078 65.078 65.078 65.078 65.078 65.078 65.078 65.078 65.078	94.330 95.999 95.125 95.941 96.455 97.049 97.816 99.115 97.287 94.367 94.274 95.053 94.021 94.763 94.7169 94.769 94.769 94.769 94.769 94.769 94.769 94.769 94.769 95.0869 97.869	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1373 1374 1376 1377 1379 1380 1383 1386 1389 1399 1399 1399 1399 1401 1403 1404 1405 1407 1409 1410 1411	O N CAB CCD NH1 C O N CAB CCD1 CCE1 CCE2 CCC OH CAB CCD1 CCE2 CCC OH CAB CCD1 CCCC OH CAB CCD1	ARG ARG ARG ARG ARG ARG ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR TYR	143 143 143 143 143 144 144 144 144 144	8.995 8.268 8.104 8.267 9.686 9.792 9.490 9.492 9.782 9.203 6.796 6.707 5.791 4.459 3.509 3.902 4.946 5.336 3.248 3.633 4.683 5.113 3.789 4.129 2.827 2.081 1.159 0.702 1.229	63.838 62.550 61.413 60.134 59.941 57.527 56.439 56.527 55.257 61.411 60.818 62.097 61.150 59.689 59.203 57.885 58.804 57.483 57.034 57.034 63.750 64.316 63.750 65.000 65.078 63.861 63.851 63.851 63.851 63.851 63.851	94.330 95.999 95.125 95.941 96.4550 97.049 97.816 99.115 97.287 94.3297 94.271 94.953 94.023 94.023 94.071 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.779 94.7799 94.779	1.00 24.93 1.00 2.00 1.00 2.86 1.00 8.64 1.00 4.03 1.00 6.91 1.00 7.13 1.00 12.97 1.00 2.00	000000000000000000000000000000000000000

		~ `	** D	146	0.2	17	66.745	97.901	1.00	2.00	0
MOTA MOTA			ILE ILE	146	0.1	68	68.278	98.048	1.00	2.00	0
ATOM	1420	CG2	ILE	146	-0.6		68.690 68.798	99.278 98.201	1.00	2.00 2.00	0
MOTA			ILE	146	1.5 1.6		70.270	98.360	1.00	2.00	ŏ
MOTA			ILE ILE	146 146	-1.1		66.143	97.851	1.00	2.00	0
MOTA MOTA		_	ILE	146	-1.6		65.927	98.881	1.00	2.00	0
MOTA		-	LYS	147	-1.6		65.839	96.668 96.594	1.00	2.00	0
MOTA			LYS	147	-3.0 -3.4		65.241 65.007	95.131		15.32	ŏ
MOTA			LYS LYS	147 147	-4.8		64.643	94.902	1.00	21.64	0
MOTA MOTA	1430		LYS	147	-5.0	24	63.248	94.280		29.48	0
ATOM	1431	CE	LYS	147	-4.7		62.128 60. 7 77	95.300 94.716		25.79 22.3 1	0
MOTA			lys Lys	147 147	-4.3 -2.8		63.928	97.345	1.00	2.00	ō
MOTA MOTA	1436 1437		LYS	147	-3.6	552	63.630	98.233	1.00	8.30	0
MOTA	1438		LEU	148	-1.8		63.180	97.023	1.00	9.36 9.36	0
MOTA	1440		LEU	148	-1.5 -0.3	582 160	61.892 61.202	97.651 97.037	1.00	2.00	ŏ
MOTA	1441		Leu Leu	148 148	-0.2	207	59.721	97.415	1.00	2.00	0
MOTA MOTA	1442 1443	CC CD1		148	-1.3	398	58.910	96.924	1.00	2.00	0
ATOM	1444	CD2	LEU	148		078	59.168	96.835 99.157	1.00	2.00 9.36	0
MOTA	1445		LEU	148	-1.4 -2.0		62.040 61.351	99.899	1.00	2.00	ŏ
MOTA	1446	0	LEU TRP	148 149	-0.		62.943	99.611	1.00	2.00	0
ATOM ATOM	1447 1449	N CA	TRP	149	-0.3	360	63.166	101.042	1.00	2.00	0
ATOM	1450	CB	TRP	149		559	64.359	101.276 102.748	1.00	9.48 13.35	Ö
MOTA	1451	CG	TRP	149		690 095	65 915	103.387	1.00	9.36	ŏ
MOTA	1452	CD2 CE2	TRP TRP	149 149		512	65.907	104.725	1.00	12.67	0
MOTA MOTA	1453 1454	CE3	TRP	149	-0.		66.946	102.951	1.00	9.36 12.84	0
MOTA	1455	CD1	TRP	149		415	64.135	103.713 104.895	1.00	14.40	ŏ
MOTA	1456	NE1	TRP	149 149		315 119	66.880	105.632	1.00	10.B2	0
MOTA	1458 1459	CZ2 CZ3	TRP TRP	149	-1.	138	67.912	103.858	`1.00	10.25	0
MOTA MOTA	1460	CH2	TRP	149	-0.			105.182 101.707	1.00	21.34	0
ATOM	1461	C	TRP	149	-1. -2.			102.647	1.00	9.36	Ō
MOTA	1462	O N	TRP LYS	1 4 9 150		429	64.434	101.216	1.00	2.00	0
MOTA MOTA	1463 1465	CA	LYS	150		756	64.777	101.725	1.00	2.00 25.82	0
ATOM	1466	CB	LYS	150		392 695	65.810 67.135	100.819	1.00	27.78	ŏ
MOTA	1467	CG	LYS	150 150		145	67.981	99.676	1.00	24.78	0
MOTA MOTA	1468 1469	CD	LYS LYS	150		236	69.442	100.053	1.00	28.33 35.49	0
ATOM	1470	NZ	LYS	150	-	243	69.673	101.132 101.786	1.00	2.00	ŏ
ATON	1474	Ç	LYS	150		661 468	63.536 63.397	102.701		28.71	0
MOTA	1475 1476	о И	LYS THR	150 151		525	62.638	100.809	1.00	2.00	0
MOTA MOTA	1478	CA	THR	151	- 5.	315	61.403	100.758	1.00	2.00 18.71	Ö
MOTA	1479	CB	THR	151	-5.	111	60.670 61.532	99.408 98.332	1.00	21.97	0
MOTA	1480		THR THR	151 151		964	59.434	99.332	1.00	19.85	0
MOTA MOTA	1482 1483	CG2 C	THR	151	-4.	920	60.487	101.925	1.00	2.00 14.71	0
ATOM	1484	Ö	THR	151		.760	59.773 60.529	102.489 102.297	1.00	2.00	0
MOTA	1485	N	PHE	152 152		.645 .161	59.729		1.00		0
MOTA	1487 1488	CA CB	PHE PHE	152	_	. 638	59.759	103.502	1.00	13.78	0
ATOM ATOM	1489	CG	PHE	152		. 956	58.586		1.00	13.78 13.78	Ö
MOTA	1490	CD1		152		.034	58.792 57.277		1.00	13.78	0
MOTA	1491	CD2		152 152		. 289 . 679	57.717	101.260	1.00	13.78	0
MOTA	1492 1493	CE1		152	-0	.648	56.194	102.578	1.00		0
MOTA MOTA	1494	CZ	PHE	152		.340	56.417 60.246		1.00		0
MOTA	1495	C	PHE	152		.767 .380	59.470	105.397	1.00		0
ATOM	1496	0	PHE	152 153		.657	61.547	7 104.944	1.00		0
MOTA MOTA	1497 1499	N CA	THR THR	153	-4	.217	62.064	106.192	1.00		0
MOTA	1500	CB	THR	153	- 4	.166	63.604	1 106.318	1.00		Ĭ

MOTA	1501	ogi 1		153	-4.912		105.265 106.293	1.00	24.34 23.93	0
MOTA	1503		THR	153	-2.739 -5.653		106.348	1.00	2.00	ŏ
MOTA	1504		THR	153 153	-6.054		107.414		29.22	0
MOTA	1505	•	CHR CB	154	-6.415		105.276	1.00	2.00	0
MOTA	1506		asp Asp	154	-7.801		105.332	1.00	2.00	0
MOTA	1508		ASP	154	-8.442	61.393	103.945		21.48	0
MOTA	1509 1510		ASP	154	-9.965		103.987	1.00		0
MOTA	1511		ASP	154	-10.576		102.971		27.37	0
MOTA	1512		ASP	154	-10.548		105.018		20.95	Ō
MOTA MOTA	1513		ASP	154	-7.889	59.806	105.801	1.00	2.00	0
ATOM	1514	-	ASP	154	-B. 78 3	59.444	106.569	1.00		0
ATOM	1515		CYS	155	-6.968		105.319	1.00	2.00	0
ATOM	1517		CYS	155	-6.951		105.691	1.00	2.00	0
ATOM	1518		CYS	155	-5.904		104.865	1.00	2.00	0
MOTA	1519		CYS	155	-5.770		105.254	1.00	2.00	0
MOTA	1520		CYS	155	-6.627		107.178	1.00	2.00	Ö
MOTA	1521		CYS	155	-7.267	56.741	107.931 107.583	1.00	8.58	ŏ
ATOM	1522		PHE	156	-5.641	58.273	107.565	1.00	8.58	Ö
MOTA	1524		PHE	156	-5.172		109.018	1.00	2.00	Ö
MOTA	1525		PHE	156	-4.056		108.437	1.00	2.00	ŏ
ATOM	1526		PHE	156	-2.766	57.585	108.206	1.00	2.00	ŏ
MOTA	1527	CD1		156	-2.537	59.847	108.160	1.00	2.00	ŏ
MOTA	1528		PHE	156	-1.756 -1. 30 9		107.710	1.00	2.00	ŏ
MOTA	1529		PHE	156	-0.518		107.662	1.00	2.00	ŏ
MOTA	1530		PHE	156 156	-0.295		107.439	1.00	2.00	0
MOTA	1531		PHE PHE	156	-6.287		109.931	1.00	8.58	0
MOTA	1532	_	PHE	156	-6.463		110.974	1.00	2.00	0
MOTA	1533 1534		asn	157	-7.055	59.758	109.572	1.00	2.00	0
MOTA	1536		ASN	157	-8.144	60.233	110.405	1.00	2.00	0
MOTA MOTA	1537		ASN	157	8.811		109.779	1.00	2.00	0
ATOM	1538		ASN	157	-7.861	62.661	109.577	1.00	2.00	0
ATOM	1539		ASN	157	-8.226	63.629	108.922	1.00	2.00	0
ATOM	1540		ASN	157	-6.669		110.143	1.00	2.00	0
MOTA	1543	С	asn	157	-9.230	59.179	110.631	1.00	2.00	Ö
MOTA	1544	0	asn	157	-10.242	59.489	111.240	1.00 1.00	2.00	Ö
MOTA	1545		CYS	158	-9.064	57.962	110.120 110.304	1.00	2.00	ŏ
MOTA	1547		CYS	158	-10.074	56.914	108.970	1.00	2.00	ŏ
MOTA	1548		CYS	158	-10.751 _. -11.898	56.567 57.864	108.370	1.00	2.00	ō
MOTA	1549	-	CYS	158	-11.656 -9.482	55.665	110.924	1.00	2.00	0
MOTA	1550	_	CYS	158 158	-10.148	54.648	111.049	1.00	2.00	0
MOTA	1551		CYS	159	-8.216		111.304	1.00	2.00	0
ATOM	1552		LEU	159	-7.433		111.945	1.00	2.00	0
MOTA	1554		LEU	159	-5.985	55.219	112.037	1.00	2.00	0
MOTA MOTA	1555 1556		LEU	159	-4.747	54.363	111.781	1.00	2.00	0
ATOM	1557	CD1		159	-4.993	53.449	110.617	1.00	2.00	0
ATOM	1558	CD2		159	-3.536	55.269	111.503	1.00	2.00	0
MOTA	1559	C	LEU	159	-8.010		113.361	1.00	2.00	0
MOTA	1560	0	LEU	159	~8. 54 8	55.367	113.980	1.00	2.00	0
ATOM	1561	N	PRO	160	-7.951	53.209	113.871	1.00	2.00 2.00	Ö
MOTA	1562	CD	PRO	160	-7.504	51.978	113.208	1.00	2.00	ŏ
MOTA	1563	CA	PRO	160	-8.473	52.896	115.209 115.269	1.00	2.00	ŏ
MOTA	1564	CB	PRO	160	-8.398	21.3/3	113.838	1.00	2.00	0
MOTA	1565	CG	PRO	160	-8.416	53.516		1.00	2.00	0
MOTA	1566	Č	PRO	160	-7.535 -6.329		116.031	1.00	2.00	0
MOTA	1567	0	PRO	160	-8.062	54.073			10.42	0
MOTA	1568	И	ILE	161 161	-7.206		118.312	1.00	10.42	0
MOTA	1570	CA	ILE	161	-7.862	55.995	118.918	1.00	2.00	0
MOTA	1571	CB	ILE	161	-8.493	56.837	117.823	1.00	2.00	0
ATOM	1572	CG2 CG1	ILE	161	-8.899	55.612	119.987	1.00	2.00	0
MOTA	1573 1574	CD1	ILE	161	-9.428	56.766	120.775	1.00	2.00	0
MOTA MOTA	1575	CDI	ILE	161	-6.741	53.859	119.492	1.00	10.42	0
ATOM	1576	Ö	ILE	161	-5.808	54.254	120.193	1.00	2.00	0
MOTA	1577	й	ALA	162	-7.387	52.712	119.720		16.67	0
ATOM	1579	CA	ALA	162	-7.036	51.832	120.837	1.00	16.67	0

ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1581 1582 1583 1585 1586 1587 1588 1589 1591 1592 1593 1594 1595 1596 1597 1598 1600 1601	CB ALA C ALA	162 163 163 163 164 164 164 165 165 166 166 166 166 167 177 177 167	-7.580
MOTA MOTA	1622 1623	OE2 GLU C GLU O GLU	167 167 167	-9.056 41.981 128.585 1.00 2.00 0 -9.634 42.138 129.657 1.00 74.79 0
MOTA	1624 1625	N LYS	168 168	-7.733 41.984 128.488 1.00 14.49 0 -6.829 42.154 129.589 1.00 14.49 0
MOTA MOTA	1627 1628	CB LYS	168	-6.956 39.798 130.600 1.00 17.49 0
MOTA MOTA	1629 1630	CC LYS	168 168	-7.677 40.404 131.802 1.00 18.27 U
MOTA	1631	CE- LYS	168 168	-9.658 40.080 133.373 1.00 29.04 0
MOTA MOTA	1632 1636	C LYS	168	-5.65 44 214 129.947 1.00 17.06 0
MOTA	1637 1638	O LYS N ILE	168 169	-5.109 43.015 128.131 1.00 2.00 0
MOTA MOTA	1640	CA ILE	169	3 151 43 267 126 648 1.00 2.00 0
MOTA MOTA	1641 1642	CB ILE	169 169	-1.989 44.184 126.302 1.00 2.00 0
ATOM	1643	CG1 ILE	169 169	-2.632 41.332 127.297 1.00 2.00 0
MOTA MOTA	1644 1645	CD1 ILE	169	-4.731 45.144 126.964 1.00 2.00 0
MOTA	1646	O ILE	169 170	-5.393 44.373 123.524 1.00 17.81 0 -4.572 46.337 127.524 1.00 17.81
MOTA MOTA	1647 1649	N PHE	170	-5.110 47.525 126.877 1.00 13.70 C
MOTA	1650	CB PHE	170 170	5 895 49.920 127.252 1.00 2.00
MOTA MOTA	1651 1652	CG PHE	170	-7.095 50.028 126.538 1.00 2.00 G
MOTA	1653	CD2 PHE		-5.092 51.036 127.976 1.00 2.00 (-7.493 51.248 125.976 1.00 2.00 (
MOTA	1654 1655	CE1 PHE		-5.473 52.282 126.836 1.00 2.00
MOTA MOTA	1656	CZ PHE	170	3 943 47.992 126.038 1.00 23.45
ATOM	1657			-2.809 47.983 126.518 1.00 2.00
MOTA	1658	, ,		

			~	101	4 200	48.375 124.790	1.00 2.00	0
MOTA	1659	N	CYS CYS	171 171	-4.208 -3.158	48.836 123.898	1.00 2.00	Ö
MOTA	1661 1662	CA CB	CYS	171	-2.952	47.843 122.766	1.00 26.14	0
ATOM ATOM	1663	SG	CYS	171	-2.524	46.203 123.305	1.00 16.13	0
ATOM	1664	c	CYS	171	-3.499	50.189 123.310	1.00 2.00	0
MOTA	1665	ŏ	CYS	171	-4.652	50.453 122.943	1.00 26.14	0
ATOM	1666	N	CYS	172	-2.495	51.058 123.265	1.00 2.00	0
ATOM	1668	CA	CYS	172	-2.635	52.378 122.668	1.00 2.00	0
ATOM	1669	CB	CYS	172	-3.403	53.356 123.597	1.00 14.54	0
ATOM	1670	SG	CYS	172	-2.593	53.995 125.094	1.00 15.67	0
MOTA	1671	С	CYS	172	-1.231	52.890 122.287	1.00 2.00 1.00 7.48	0
MOTA	1672	0	CYS	172	-0.227	52.277 122.657 53.951 121.488	1.00 2.00	0
MOTA	1673	N	HIS	173	-1.154	53.951 121.488 54.461 121.112	1.00 2.00	ŏ
MOTA	1675	CA	HIS	173	0.146 0.815	55.130 122.297	1.00 2.00	ŏ
ATOM	1676	C	HIS HIS	173 173	1.893	54.707 122.722	1.00 2.00	ŏ
MOTA	1677	O CB	HIS	173	0.056	55.457 119.944	1.00 2.00	0
MOTA	1678 1679	CG	HIS	173	1.377	56.074 119.588	1.00 2.00	0
ATOM ATOM	1680		HIS	173	2.487	55.347 119.234	1.00 2.00	0
ATOM	1682		HIS	173 -	1.774	57.372 119.585	1.00 2.00	0
MOTA	1683		HIS	173	3.123	57.441 119.241	1.00 2.00	0
ATOM	1684		HIS	173	3.492	56.189 119.045	1.00 2.00	0
MOTA	1685	N	GLY	174	0.168	56.172 122.817	1.00 2.00	0
MOTA	1687	CA	GLY	174	0.711	56.933 123.935	1.00 2.00	0
ATOM	1688	С	GLY	174	0.568	56.450 125.386	1.00 2.00 1.00 2.00	0
MOTA	1689	0	GLY	174	1.556	56.111 126.041 56.453 125.913	1.00 2.00 1.00 16.98	Ö
MOTA	1690	N	GLY	175	-0.648 -0.830	56.453 125.913 56.028 127.283	1.00 16.98	ŏ
MOTA	1692	CA	GLY	175	-2.227	56.343 127.763	1.00 16.98	ŏ
ATOM	1693	C	GLY	175 175	-3.190	56.231 126.999	1.00 66.77	Ō
ATOM	1694 1695	O N	GLY LEU	176	-2.341	56.764 129.021	1.00 11.28	0
ATOM ATOM	1697	N CA	LEU	176	-3.637	57.063 129.604	1.00 11.28	0
ATOM	1698	CB	LEU	176	-3.740	56.427 130.976	1.00 2.00	0
ATOM	1699	CG	LEU	176	-3.443	54.934 130.966	1.00 2.00	0
ATOM	1700	CD1	LEU	176	-3.469	54.407 132.374	1.00 2.00	0
ATOM	1701	CD2	LEU	176	-4.463	54.218 130.121	1.00 2.00	0
MOTA	1702	С	LEU	176	-3.876	58.545 129.692	1.00 11.28 1.00 2.00	0
MOTA	1703	0	LEU	176	-2.943	59.329 129.645 58.913 129.850	1.00 2.00 1.00 6.18	Ö
MOTA	1704	N	SER	177	-5.138	58.913 129.850 60.304 129.918	1.00 6.86	ŏ
MOTA	1706	CA	SER	177	-5. 567 -6. 47 6	60.587 128.718	1.00 12.19	ŏ
ATOM	1707	CB	SER	177 177	-7.189	61.804 128.835	1.00 12.19	0
ATOM	1708	og	SER	177	-6.356	60.573 131.193	1.00 7.79	0
MOTA	1710 1711	CO	SER	177	-7.170	59.749 131.606	1.00 12.19	0
ATOM ATOM	1712	N	PRO	178	-6.150	61.736 131.824	1.00 2.00	0
ATOM	1713	င်ာ	PRO	178	-5.223	62.827 131.505	1.00 29.89	0
ATOM	1714	CA	PRO	178	-6.895	62.054 133.041	1.00 2.00	0
MOTA	1715	CB	PRO	178	-6.231	63.337 133.518	1.00 25.57	0
MOTA	1716	CG	PRO	178	-5.842	63.973 132.274	1.00 21.36	0
MOTA	1717	C	PRO	178	-8.394	62.266 132.757	1.00 2.00	
MOTA	1718		200	178				
MOTA		0	PRO		-9.140	62.740 133.617	1.00 24.04	0
	1719	N	ASP	179	-8.821	61.933 131.543	1.00 38.94	0
ATOM	1719 1721	N CA	ASP ASP	179 179	-8.821 -10.206	61.933 131.543 62.078 131.129	1.00 38.94 1.00 37.05	
MOTA MOTA	1719 1721 1722	N CA CB	ASP ASP ASP	179 179 179	-8.821 -10.206 -10.264	61.933 131.543 62.078 131.129 62.870 129.837	1.00 38.94 1.00 37.05 1.00 33.34	0
MOTA MOTA MOTA	1719 1721 1722 1723	N CA CB CG	ASP ASP ASP	179 179 179 179	-8.821 -10.206 -10.264 -9.964	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71	0 0 0 0
MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724	N CA CB CG OD1	ASP ASP ASP ASP	179 179 179 179 179	-8.821 -10.206 -10.264	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17	00000
ATOM MOTA ATOM ATOM ATOM MOTA	1719 1721 1722 1723 1724 1725	N CA CB CG OD1 OD2	ASP ASP ASP ASP ASP	179 179 179 179	-8.821 -10.206 -10.264 -9.964 -10.923	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.0386 64.728 129.889 60.735 130.923	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05	000000
MOTA MOTA MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724 1725 1726	N CA CB CG OD1 OD2 C	ASP ASP ASP ASP	179 179 179 179 179 179	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.889 60.735 130.923 60.638 130.873	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46	0000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724 1725 1726 1727	N CA CB CG OD1 OD2	ASP ASP ASP ASP ASP ASP	179 179 179 179 179 179	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.386 60.735 130.923 60.638 130.873 59.699 130.791	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 15.43	00000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724 1725 1726	N CA CB CG OD1 OD2 C	ASP ASP ASP ASP ASP ASP ASP	179 179 179 179 179 179 179 179 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.057 -10.570	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.836 60.735 130.923 60.638 130.873 59.699 130.791 58.362 130.590	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 15.43 1.00 12.98	000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724 1725 1726 1727 1728 1730 1731	N CA CB CG OD1 OD2 C	ASP ASP ASP ASP ASP ASP LEU LEU	179 179 179 179 179 179 179 179 180 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.057 -10.570 -9.446	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.836 60.735 130.923 60.638 130.873 59.699 130.791 58.362 130.590 57.467 130.058	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 15.43 1.00 12.98 1.00 2.00	00000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724 1725 1726 1727 1728 1730 1731 1732	N CB CG OD1 OD2 C O N CA CB	ASP ASP ASP ASP ASP ASP LEU LEU LEU	179 179 179 179 179 179 179 179 180 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.570 -9.446 -9.867	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.889 60.735 130.923 60.638 130.873 59.699 130.791 58.362 130.590 57.467 130.058 56.186 129.338	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 12.98 1.00 2.00 1.00 2.00	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1719 1721 1722 1723 1724 1725 1726 1727 1728 1730 1731 1732 1733	N CA CB CG OD1 OD2 C O N CA CB CG CD1	ASP ASP ASP ASP ASP ASP LEU LEU LEU LEU	179 179 179 179 179 179 179 180 180 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.570 -9.446 -9.867 -10.706	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.889 60.735 130.923 60.638 130.923 60.638 130.791 59.699 130.791 58.362 130.590 57.467 130.058 56.186 129.338 56.511 128.110	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 15.43 1.00 12.00 1.00 2.00 1.00 2.00	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1719 1721 1722 1723 1724 1725 1726 1727 1728 1730 1731 1732 1733 1734	N CA CB CG OD1 OD2 C O N CA CB CG CD1 CD2	ASP ASP ASP ASP ASP ASP LEU LEU LEU LEU LEU	179 179 179 179 179 179 179 180 180 180 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.057 -10.570 -9.446 -9.867 -10.706 -8.633	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.386 65.045 130.386 64.728 129.889 60.735 130.923 60.638 130.873 59.699 130.791 58.362 130.590 57.467 130.595 56.186 129.338 56.511 128.110 55.413 128.953	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 15.43 1.00 12.00 1.00 2.00 1.00 2.00 1.00 2.00	00000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	1719 1721 1722 1723 1724 1725 1726 1727 1728 1730 1731 1732 1733 1734 1735	N CA CB CG OD1 CD2 C C CD1 CD2 C	ASP ASP ASP ASP ASP ASP LEU LEU LEU LEU LEU	179 179 179 179 179 179 179 180 180 180 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.057 -10.570 -9.446 -9.867 -10.706 -8.633 -11.157	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.048 65.045 130.386 64.728 129.889 60.735 130.923 60.638 130.873 59.699 130.791 58.362 130.590 57.467 130.590 57.467 130.590 556.186 129.338 56.511 128.110 55.413 128.953 57.806 131.902	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 40.05 1.00 34.46 1.00 15.43 1.00 12.98 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.43	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1719 1721 1722 1723 1724 1725 1726 1727 1728 1730 1731 1732 1733 1734	N CA CB CG OD1 OD2 C O N CA CB CG CD1 CD2	ASP ASP ASP ASP ASP ASP LEU LEU LEU LEU LEU	179 179 179 179 179 179 179 180 180 180 180	-8.821 -10.206 -10.264 -9.964 -10.923 -8.786 -10.871 -12.096 -10.057 -10.570 -9.446 -9.867 -10.706 -8.633	61.933 131.543 62.078 131.129 62.870 129.837 64.320 130.386 65.045 130.386 64.728 129.889 60.735 130.923 60.638 130.873 59.699 130.791 58.362 130.590 57.467 130.595 56.186 129.338 56.511 128.110 55.413 128.953	1.00 38.94 1.00 37.05 1.00 33.34 1.00 30.55 1.00 31.71 1.00 32.17 1.00 40.05 1.00 34.46 1.00 15.43 1.00 12.00 1.00 2.00 1.00 2.00 1.00 2.00	00000000000000

						56 017	*** 002	1.00 5.88	0
MOTA	1739	Cy	GLN	181	-13.184		132.992 133.375	1.00 5.88 1.00 32.99	ŏ
MOTA	1740	CB	GLN	181	-14.379 -14.002	•	133.900	1.00 36.07	ŏ
MOTA	1741	CC	GLN	181	-15.101		133.676	1.00 38.35	ŏ
ATOM	1742	CD	GLN	181	-16.285		133.877	1.00 41.53	ŏ
ATOM	1743		GLN	181 181	-14.717		133.242	1.00 38.39	ō
MOTA	1744	NE2	GLN	181	-13.672		132.500	1.00 5.88	ō
MOTA	1747	C	GLN	181	-13.572		133.205	1.00 24.31	ō
MOTA	1748	0	GLN SBR	182	-14.188		131.274	1.00 2.20	0
ATOM	1749	N	SER	182	-14.651		130.664	1.00 9.71	Ŏ
MOTA	1751	CA	SER	182	-16.174		130.614	1.00 15.80	Ó
MOTA	1752	CB OG	SER	182	-16.668		129.433	1.00 17.25	0
MOTA	1753		SER	182	-14.112		129.248	1.00 3.77	0
MOTA	1755	C	SER	182	-13.359		128.751	1.00 15.80	Ó
MOTA	1756	O N	MET	183	-14.526		128.615	1.00 2.00	0
MOTA	1757		MET	183	-14.156		127.251	1.00 2.00	0
MOTA	1759	CA CB	MET	183	-14.100		127.064	1.00 2.00	0
MOTA	1760	CG	MET	183	-13.171		128.048	1.00 2.00	0
MOTA	1761 1762	SD	MBT	183	-11.620		128.022	1.00 2.00	0
ATOM	1763	CE	MET	183	-10.520		127.900	1.00 2.00	0
MOTA		C	MET	183	-15.204		126.326	1.00 2.00	0
MOTA	1764		MET	183	-14.959		125.129	1.00 2.00	0
ATOM	1765	0	GLU	184	-16.370		126.882	1.00 2.00	0
ATOM	1766	N		184	-17.432		126.082	1.00 2.00	0
MOTA	1768	CA	GLU	184	-18.668		126.910	1.00 6.25	0
MOTA	1769	CB	GLU	184	-19.830		126.073	1.00 8.82	0
ATOM	1770	CG	GLU		-20.273		125.068	1.00 13.01	0
MOTA	1771	CD	GLU	184	-19.846		125.181	1.00 14.09	0
MOTA	1772	OE1		184	-21.064		124.164	1.00 20.29	0
MOTA	1773	OE2	GLU	184	-16.978	55.586	125.481	1.00 2.00	0
MOTA	1774	C	GLU	184	-17.399		124.392	1.00 23.33	0
MOTA	1775	0	GLU	184	-16.117		126.198	1.00 12.32	0
MOTA	1776	N	GLN	185	-15.599		125.714	1.00 14.30	0
MOTA	1778	CA	GLN	185	-14.697		126.758	1.00 43.49	0
MOTA	1779	CB	GLN	185	-15.454	58.629	127.990	1.00 47.40	0
MOTA	1780	cc	GLN	185	-14.537		129.139	1.00 49.32	Ō
MOTA	1781	CD	GLN	185	-13.994		129.753	1.00 56.49	O
ATOM	1782	OE1	GLN	185	-14.350	60.208	129.437	1.00 49.76	0
MOTA	1783	NE2	GLN	185 185	-14.834	57.307	124.432	1.00 16.93	0
ATOM	1786	C	GLN	185	-14.973		123.461	1.00 43.79	0
MOTA	1787	0	GLN	186	-14.044		124.420	1.00 5.67	0
MOTA	1788	N	ILE	186	-13.280		123.235	1.00 5.67	0
MOTA	1790	CA		186	-12.436	54.613	123.470	1.00 16.18	0
MOTA	1791	CB	ILE	186	-11.675	54.243	122.208	1.00 11.85	0
MOTA	1792	CG2	ILE	186	-11.489	54.825	124.642	1.00 13.18	0
MOTA	1793	CG1		186	-10.545	55.954	124.459	1.00 17.92	0
MOTA	1794	CD1		186	-14.294	55.566	122.143	1.00 5.67	0
MOTA	1795	C	ILE	186	-14.260	56.145	121.060	1.00 19.35	0
MOTA	1796	0	ILE	187	-15.207	54.655	122.471	1.00 17.59	0
MOTA	1797	N	ARG	187	-16.243		121.561	1.00 16.33	0
ATOM	1799	CA	ARG	187	-17.141	53.165	122.237	1.00 20.41	0
MOTA	1800	CB	ARG	187	-16.468	51.900	122.723	1.00 29.32	0
MOTA	1801	CG	ARG ARG	187	-17.497	51.006	123.447	1.00 31.43	0
MOTA	1802	CD	ARG	187	-16.888	49.888	124.165	1.00 38.34	0
ATOM	1803	NE	ARG	187	-16.311	48.838	123.580	1.00 34.30	0
ATOM	1805	CZ		187	-16.256	48.747	122.252	1.00 41.37	0
ATOM	1806	NH1		187	-15.783	47.873	124.322	1.00 34.50	0
MOTA	1809	NH2	ARG ARG	187	-17.148	55.302	121.011	1.00 15.89	0
MOTA	1812	C	ARG	187	-17.937	55.032	120.108	1.00 20.82	0
MOTA	1813	0		188	-17.071	56.524	121.529	1.00 2.00	0
MOTA	1814	N	ARG	188	-17.964	57.557	121.012	1.00 2.00	0
ATOM	1816	CA	ARG ARG	188	-18.878	58.105	122.106	1.00 31.38	0
MOTA	1817	CB	ARG	188	-18.184	58.827	123.228	1.00 29.44	0
MOTA	1818	CG	מאת		10.202	59.586	124.034	1.00 31.32	0
MOTA			APC	188	-19.202	33.300	124.034		
	1819	CD	ARG	188 188	-19.202 -20.410		124.237	1.00 33.18	0
MOTA	1819 1820	CD NE	ARG	188	-20.410	58.802 59.285	124.237		0
MOTA MOTA	1819 1820 1822	CD NE CZ	ARG ARG	188 188	-20.410 -21.637	58.802 59.285	124.237	1.00 33.18	
MOTA	1819 1820	CD NE CZ NH	ARG	188	-20.410	58.802	124.237 124.112	1.00 33.18 1.00 29.63	0



ATOM 1845 O ARG 188 -17.758	MOTA	1829	С	ARG	188	-17.255	58.682	120.332	1.00 2.00	0
ATOM 1831 CA LIE 189 -15.263 59.333 119.336 1.00 16.69 O ATOM 1835 CB LIE 189 -13.759 59.043 119.345 1.00 16.67 O ATOM 1836 CB LIE 189 -13.759 59.043 119.345 1.00 17.65 O ATOM 1837 CD LIE 189 -13.759 59.043 119.345 1.00 17.65 O ATOM 1837 CD LIE 189 -13.61 59.205 120.826 1.00 17.65 O ATOM 1837 CD LIE 189 -13.61 59.205 120.826 1.00 17.65 O ATOM 1839 C LIE 189 -13.61 59.205 120.826 1.00 17.65 O ATOM 1839 C LIE 189 -15.620 59.244 117.656 1.00 19.51 O ATOM 1839 C LIE 189 -15.620 59.244 117.656 1.00 19.51 O ATOM 1840 N MET 190 -16.514 60.420 115.685 1.00 22.12 O ATOM 1841 CB MET 190 -16.514 60.420 115.685 1.00 24.12 O ATOM 1843 CB MET 190 -17.951 61.812 115.320 1.00 46.39 O ATOM 1845 SD MET 190 -17.950 61.977 113.826 1.00 46.39 O ATOM 1846 CE MET 190 -17.950 61.977 113.826 1.00 46.39 O ATOM 1846 CE MET 190 -15.673 60.194 115.320 1.00 46.39 O ATOM 1846 CE MET 190 -15.673 60.194 115.00 50.365 10.00 50.38 O ATOM 1846 CE MET 190 -15.673 60.194 115.105 1.00 58.03 O ATOM 1846 CE MET 190 -15.73 60.194 115.105 1.00 52.53 6 O ATOM 1851 CA ARG 191 -13.732 58.738 113.731 1.00 2.00 O ATOM 1852 CB ARG 191 -13.732 58.738 113.772 1.00 2.00 O ATOM 1855 CB ARG 191 -13.732 58.738 113.772 1.00 1.131 O ATOM 1855 CB ARG 191 -13.732 58.738 113.762 1.00 11.31 O ATOM 1855 CB ARG 191 -13.734 55.366 113.416 1.00 2.00 O ATOM 1855 CB ARG 191 -13.735 59.904 113.752 1.00 11.31 O ATOM 1856 CD ARG 191 -13.744 55.663 113.416 1.00 2.00 O ATOM 1857 CZ ARG 191 -13.743 59.35 191.3752 1.00 11.31 O ATOM 1858 NA ARG 191 -13.744 55.663 113.416 1.00 2.00 O ATOM 1856 CD ARG 191 -13.753 60.064 113.407 1.00 2.00 O ATOM 1857 CZ ARG 191 -13.754 50.064 113.607 10.00 10.00 O ATOM 1858 NA ARG 191 -13.754 50.064 113.607 10.00 10.00 O ATOM 1858 NA ARG 191 -13.754 50.064 113.607 10.00 10.00 O ATOM 1858 NA ARG 191 -13.754 50.00 NA O ATOM 1868 CB PRO 192 -14.743 59.73 113.607 1.00 2.00 O ATOM 1868 CB PRO 192 -14.743 59.73 113.607 1.00 2.00 O ATOM 1868 CB PRO 192 -14.743 59.73 113.607 1.00 2.00 O ATOM 1879 CG PRO 196 -6.675 60.677 113.207 1.00 2.00 O ATOM 1889 CB PRO 192				ARG	188	-17.758	59.804	120.250	1.00 31.89	0
ATOM 1834 CB ILE 189 -13.759 59.043 119.345 1.00 16.67 0 ATOM 1835 CG2 ILE 189 -13.411 59.205 120.826 1.00 17.65 0 ATOM 1836 CG1 ILE 189 -13.411 59.205 120.826 1.00 17.65 0 ATOM 1837 CD1 ILE 189 -13.411 59.205 120.826 1.00 17.65 0 ATOM 1838 C ILE 189 -15.620 59.244 117.656 1.00 19.51 0 ATOM 1838 C ILE 189 -15.620 59.244 117.656 1.00 19.51 0 ATOM 1840 N MET 190 -16.588 60.342 117.108 1.00 22.12 0 ATOM 1842 CA MET 190 -16.518 60.342 117.108 1.00 22.12 0 ATOM 1843 CB MET 190 -17.051 61.812 115.820 1.00 46.39 0 ATOM 1844 CG MET 190 -17.056 61.871 113.826 1.00 46.39 0 ATOM 1845 SD MET 190 -17.206 61.812 115.820 1.00 46.39 0 ATOM 1846 CE MET 190 -17.206 61.971 113.826 1.00 22.38 0 ATOM 1846 CE MET 190 -17.820 61.558 113.312 1.00 58.03 0 ATOM 1847 C MET 190 -15.173 66.158 113.312 1.00 58.03 0 ATOM 1848 C MET 190 -15.173 66.158 113.312 1.00 52.38 0 ATOM 1848 C MET 190 -15.173 66.158 113.312 1.00 25.36 0 ATOM 1847 C MET 190 -15.173 66.158 113.312 1.00 25.36 0 ATOM 1858 C MAR 191 -13.731 58.738 113.772 1.00 2.00 0 ATOM 1851 CA ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATOM 1852 CB ARG 191 -13.731 58.738 113.772 1.00 2.00 0 ATOM 1853 CC ARG 191 -13.731 55.738 113.747 1.00 2.00 0 ATOM 1854 CD ARG 191 -13.731 55.693 113.461 1.00 2.00 0 ATOM 1855 CA ARG 191 -13.139 54.935 113.747 1.00 2.00 0 ATOM 1866 CD ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1867 CZ ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1867 CC ARG 191 -13.124 55.665 113.670 1.00 2.00 0 ATOM 1867 CC ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1867 C ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1867 C ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1867 C ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1867 C ARG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1868 C A RG 191 -13.124 55.665 113.467 1.00 2.00 0 ATOM 1869 C B RO 192 -14.626 60.647 113.202 1.00 60.00 0 ATOM 1869 C B RO 192 -14.626 60.647 113.202 1.00 60.00 0 ATOM 1869 C B RO 192 -14.626 60.647 113.00 11.00 2.00 0 ATOM 1869 C B RO 192 -14.626 60.647 113.00 11.00 2.00 0 ATOM 1869 C B RO 192							59.333	119.136		
NATION 1835 CG2 LLE 189							59.043	119.345		
ATOM 1837 CD1 TLE 18511.970 58.946 121.773 1.00 16.74 0 ATOM 1838 C ILE 18915.620 59.244 117.656 1.00 19.51 0 ATOM 1839 O ILE 18915.448 58.184 117.044 1.00 25.93 0 ATOM 1809 O ILE 18915.448 58.184 117.044 1.00 25.93 0 ATOM 1804 CA MET 19016.514 60.420 115.685 1.00 24.92 0 ATOM 1842 CA MET 19016.514 60.420 115.685 1.00 24.92 0 ATOM 1842 CB MET 19017.366 61.977 113.826 1.00 46.97 0 ATOM 1844 CG MET 19017.366 61.977 113.826 1.00 46.97 0 ATOM 1845 SD MET 19017.366 61.977 113.826 1.00 52.38 0 ATOM 1846 CB MET 19016.271 64.266 112.598 1.00 52.38 0 ATOM 1848 O MET 19016.271 64.266 112.598 1.00 52.38 0 ATOM 1848 O MET 19015.173 60.194 115.015 1.00 25.36 0 ATOM 1849 N ARG 19115.023 59.057 114.346 1.00 2.00 0 ATOM 1851 CA ARG 19113.732 58.738 113.772 1.00 2.00 0 ATOM 1852 CB ARG 19113.732 58.738 113.772 1.00 1.031 0 ATOM 1854 CD ARG 19113.139 55.799 113.752 1.00 11.31 0 ATOM 1855 NE ARG 19113.139 55.799 113.752 1.00 11.31 0 ATOM 1856 CD ARG 19113.139 55.799 113.752 1.00 11.31 0 ATOM 1858 NNI ARG 19113.139 55.799 113.752 1.00 10.31 0 ATOM 1856 NE ARG 19113.139 55.799 113.752 1.00 10.31 0 ATOM 1866 C ARG 19113.139 55.799 113.752 1.00 10.31 0 ATOM 1866 N PRO 19213.685 60.065 111.670 1.00 2.00 0 ATOM 1866 N PRO 19214.926 52.998 113.852 1.00 62.00 0 ATOM 1867 CC PRO 19214.926 52.999 113.752 1.00 12.00 0 ATOM 1868 CA PRO 19214.926 52.999 113.752 1.00 12.00 0 ATOM 1868 CA PRO 19214.926 52.999 113.752 1.00 12.00 0 ATOM 1868 CA PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1868 CA PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1868 CA PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1870 CG PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1870 CG PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1870 CG PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1870 CG PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1870 CG PRO 19214.926 52.999 113.752 1.00 0.00 0 ATOM 1880 C ARG 19115.995 60.267 113.202 1.00 6.00 0 ATOM 1880 C ARG 19115.995 60.267							59.973	118.493		
ATOM 1838 C			-				58.946	121.173		
ATOM 1840 N MET 190 -15.848 58.184 117.044 1.00 22.93 0 ATOM 1840 N MET 190 -16.518 60.342 117.104 1.00 22.11 0 ATOM 1843 CB MET 190 -15.515 60.342 117.044 1.00 22.12 0 ATOM 1843 CB MET 190 -17.366 60.420 115.685 1.00 24.92 0 ATOM 1844 CG MET 190 -17.366 61.977 113.826 1.00 46.39 0 ATOM 1845 SD MET 190 -17.366 61.977 113.826 1.00 46.39 0 ATOM 1846 CE MET 190 -17.366 61.977 113.825 1.00 52.38 0 ATOM 1846 CE MET 190 -16.271 64.266 112.598 1.00 52.38 0 ATOM 1846 CE MET 190 -16.271 64.266 112.598 1.00 52.38 0 ATOM 1848 O MET 190 -14.287 61.041 115.106 1.00 32.576 0 ATOM 1849 N ARG 191 -15.033 59.057 114.346 1.00 23.56 0 ATOM 1851 CA ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATOM 1852 CB ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1854 CG ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1855 NE ARG 191 -11.788 55.799 113.752 1.00 11.31 0 ATOM 1856 NE ARG 191 -13.131 54.935 113.762 1.00 2.00 0 ATOM 1858 NHI ARG 191 -11.312 53.663 113.416 1.00 2.00 0 ATOM 1858 NHI ARG 191 -11.978 55.799 113.752 1.00 11.31 0 ATOM 1856 N PRO 192 -11.995 50.085 113.047 1.00 2.00 0 ATOM 1866 N PRO 192 -11.995 60.067 113.007 1.00 2.00 0 ATOM 1866 N PRO 192 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1866 N PRO 192 -11.995 60.067 113.007 1.00 2.00 0 ATOM 1867 CD PRO 192 -14.701 55.900 111.670 1.00 37.06 0 ATOM 1868 CA PRO 192 -11.995 60.067 113.007 1.00 2.00 0 ATOM 1868 CA PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.701 55.900 111.007 1.00 2.00 0 ATOM 1870 CG PRO 192 -17.000 111.100 1.00 2.00 0 ATOM 1870 CG PRO 192 -17.000 1.000							59.244	117.656		
ATOM 1840 N MET 190 -16.158 60.342 117.108 1.00 24.12 0 ATOM 1842 CA MET 190 -17.051 61.812 115.325 1.00 46.39 0 ATOM 1843 CB MET 190 -17.051 61.812 115.325 1.00 46.39 0 ATOM 1844 CG MET 190 -17.051 61.812 115.325 1.00 58.03 0 ATOM 1845 SD MET 190 -17.820 63.658 113.312 1.00 58.03 0 ATOM 1846 CE MET 190 -17.820 63.658 113.312 1.00 58.03 0 ATOM 1846 C MET 190 -15.173 60.194 115.015 1.00 25.36 0 ATOM 1847 C MET 190 -15.173 60.194 115.015 1.00 25.36 0 ATOM 1848 O MET 190 -15.173 60.194 115.015 1.00 25.36 0 ATOM 1851 CA ARG 191 -13.731 57.384 113.772 1.00 2.00 0 ATOM 1852 CB ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1853 CG ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1854 CD ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1855 NE ARG 191 -13.139 54.935 113.752 1.00 11.31 0 ATOM 1855 NE ARG 191 -13.139 54.935 113.752 1.00 11.31 0 ATOM 1857 CZ ARG 191 -13.139 54.935 113.752 1.00 11.31 0 ATOM 1858 NH1 ARG 191 -13.955 3.085 113.416 1.00 2.00 0 ATOM 1864 C ARG 191 -13.108 53.085 113.407 1.00 2.00 0 ATOM 1857 CZ ARG 191 -13.108 53.085 113.407 1.00 2.00 0 ATOM 1865 NH ARG 191 -13.108 53.085 113.407 1.00 2.00 0 ATOM 1866 N PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1867 CD PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1868 CA PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1868 CD ARG 191 -13.95 59.769 112.857 1.00 2.00 0 ATOM 1868 CD ARG 191 -13.084 59.769 112.857 1.00 2.00 0 ATOM 1868 CD ARG 191 -13.139 54.93 113.752 1.00 11.31 0 ATOM 1868 CD ARG 191 -13.139 54.93 113.752 1.00 11.31 0 ATOM 1868 CD ARG 191 -13.139 54.93 113.752 1.00 11.31 0 ATOM 1868 CD ARG 191 -13.139 55.799 113.357 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -13.64 59.769 112.857 1.00 2.00 0 ATOM 1862 CD ARG 191 -13.139 54.93 113.752 1.00 0.0 0.0 0 ATOM 1863 NH1 ARG 191 -13.93 1.00 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.						-15.448	58.184	117.044		
ATOM 1844 CG HET 190 -17.361 61.812 115.320 1.00 46.39 0 ATOM 1844 CG HET 190 -17.366 61.977 113.826 1.00 46.97 0 ATOM 1845 SD HET 190 -17.220 63.658 113.312 1.00 58.03 0 ATOM 1846 CE HET 190 -16.271 64.265 112.598 1.00 52.38 0 ATOM 1846 CE HET 190 -16.271 64.265 112.598 1.00 52.38 0 ATOM 1848 O MET 190 -15.373 60.194 115.015 1.00 25.36 0 ATOM 1849 N ARG 191 -15.023 59.057 114.346 1.00 2.50 0 ATOM 1851 CA ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATOM 1852 CB ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1853 CG ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1854 CD ARG 191 -13.732 55.799 113.752 1.00 11.31 0 ATOM 1855 NE ARG 191 -13.139 54.935 113.762 1.00 2.00 0 ATOM 1856 NH ARG 191 -13.139 54.935 113.762 1.00 2.00 0 ATOM 1858 NH ARG 191 -11.978 55.799 113.752 1.00 12.31 0 ATOM 1858 NH ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1858 NH ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1866 C ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1866 N PEO 192 -11.985 60.065 111.670 1.00 37.066 0 ATOM 1866 N PEO 192 -11.985 60.065 111.670 1.00 37.06 0 ATOM 1866 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1866 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1867 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1868 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1868 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1869 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1870 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1880 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1881 O THR 193 -11.995 60.267 113.207 1.00 2.00 0 ATOM 1870 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1880 C PEO 192 -11.085 60.065 111.670 1.00 37.06 0 ATOM 1880 C PEO 192 -14.73 59.973 111.09 60.00 1.00 0.00 0 ATOM 1880 C PEO 192 -14.73 59.973 111.09 60.00 0.00 0 ATOM 1880 C PEO 192 -14.73 59.973 111.00 0.00 0.00 0.00 0.00 0.00 0.00 0		1840					60.342	117.108		
ATCH 1844 CG HET 190 -17.366 61.977 113.826 1.00 46.97 0 ATCH 1845 SD HET 190 -17.820 63.658 113.312 1.00 58.03 0 ATCH 1846 CE HET 190 -16.271 64.265 112.598 1.00 52.38 0 ATCH 1848 C HET 190 -15.173 60.194 115.015 1.00 25.36 0 ATCH 1848 C HET 190 -15.173 60.194 115.015 1.00 25.36 0 ATCH 1848 C HET 190 -15.173 60.194 115.015 1.00 25.36 0 ATCH 1848 C HET 190 -14.287 61.041 115.106 1.00 32.57 0 ATCH 1849 N ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATCH 1851 CA ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATCH 1852 CG ARG 191 -12.300 56.900 112.821 1.00 11.31 0 ATCH 1853 CG ARG 191 -11.978 55.799 113.752 1.00 11.31 0 ATCH 1855 NE ARG 191 -13.19 54.935 113.762 1.00 2.00 0 ATCH 1855 NE ARG 191 -13.19 54.935 113.762 1.00 2.00 0 ATCH 1857 CZ ARG 191 -13.124 53.663 113.461 1.00 2.00 0 ATCH 1865 NE ARG 191 -13.19 54.935 113.072 1.00 2.00 0 ATCH 1866 N PRO 191 -13.19 54.935 113.072 1.00 2.00 0 ATCH 1867 CD PRO 192 -13.685 60.267 113.202 1.00 6.10 0 ATCH 1867 CD PRO 192 -13.685 60.267 113.202 1.00 6.10 0 ATCH 1867 CD PRO 192 -13.685 60.267 113.202 1.00 6.10 0 ATCH 1867 CD PRO 192 -14.414 61.335 109.749 1.00 2.00 0 ATCH 1873 N THR 193 -11.527 62.348 112.683 1.00 2.00 0 ATCH 1873 N THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1875 CA THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1875 CA THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1875 CA THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1875 CA THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1870 CG THR 193 -11.527 62.348 113.454 1.00 2.00 0 ATCH 1880 C A SP 194 -9.394 64.879 113.572 1.00 12.00 0 ATCH 1880 C A SP 194 -9.394 64.879 113.572 1.00 12.00 0 ATCH 1880 C B THR 193 -11.526 63.063 111.673 1.00 2.00 0 ATCH 1880 C B THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1880 C B THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1880 C B THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1880 C B THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1880 C B THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1880 C B THR 193 -11.527 62.348 112.168 1.00 2.00 0 ATCH 1880 C B THR 193 -11.				_			61.812	115.320		
ATOM 1846 CE MET 190 -17.8200 63.658 113.311 1.00 38.07 0 ATOM 1847 C MET 190 -15.271 64.266 112.598 1.00 52.386 0 ATOM 1848 O MET 190 -15.173 60.194 115.015 1.00 25.36 0 ATOM 1848 O MET 190 -14.287 61.041 115.106 1.00 32.57 0 ATOM 1849 N ARG 191 -13.731 57.384 113.346 1.00 2.00 0 ATOM 1851 CA ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1852 CB ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1853 CG ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1854 CD ARG 191 -11.978 55.799 113.752 1.00 11.31 0 ATOM 1855 NE ARG 191 -11.1998 55.799 113.752 1.00 11.31 0 ATOM 1857 CZ ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1858 NH1 ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1866 C ARG 191 -11.995 50.067 112.857 1.00 2.00 0 ATOM 1866 N PRO 192 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1866 N PRO 192 -14.915 59.769 112.857 1.00 2.00 0 ATOM 1866 N PRO 192 -13.685 60.065 111.670 1.00 37.066 0 ATOM 1866 CA PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1867 CD PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1869 CB PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1871 C PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 112.188 1.00 2.00 0 ATOM 1876 CB THR 193 -11.627 63.063 114.540 1.00 2.00 0 ATOM 1877 CG2 THR 193 -11.627 63.063 113.591 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.592 1.00 31.66 0 ATOM 1880 C THR 193 -11.627 63.063 113.591 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.509 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.099 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 114.500 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 114.500 9.00 37.24 0 ATOM 1880 C THR 193 -11.627 63.063 114.500 9.00 37.24 0 ATOM 1880 C THR 193 -11.627 63.063 114.500 9.00 37.24 0 ATOM 1880 C THR 193 -10.666 63.061 113.522 1.00 2.00 0 ATOM 1880 C THR 193 -10.666 63.061 113.522 1.00 2.00 0 ATOM 1880 C THR 193 -1							61.977	113.826		
ATOM 1847 C MET 190			SD				63.658	113.312		
ATOM 1848 0 MET 190 -14.287 61.041 115.106 1.00 32.57 0 ATOM 1849 N ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATOM 1851 CA ARG 191 -13.732 58.738 113.772 1.00 2.00 0 ATOM 1852 CB ARG 191 -12.300 56.900 112.821 1.00 11.31 0 ATOM 1853 CG ARG 191 -12.300 56.900 112.821 1.00 11.31 0 ATOM 1855 NE ARG 191 -11.978 55.799 113.752 1.00 2.00 0 ATOM 1855 NE ARG 191 -11.978 55.799 113.752 1.00 2.00 0 ATOM 1855 NE ARG 191 -13.139 54.935 113.762 1.00 2.00 0 ATOM 1857 NE ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1858 NH1 ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1864 C ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1865 NH2 ARG 191 -11.985 60.267 113.202 1.00 6.10 0 ATOM 1866 N PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1865 C ARG 191 -13.084 59.769 112.857 1.00 2.00 0 ATOM 1865 C ARG 191 -13.995 60.267 113.202 1.00 6.10 0 ATOM 1865 C ARG 191 -13.995 60.267 113.202 1.00 6.10 0 ATOM 1865 C RO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1870 RO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1870 RO 192 -14.114 61.335 109.749 1.00 2.00 0 ATOM 1870 RO 192 -14.114 61.335 109.749 1.00 2.00 0 ATOM 1870 RO 192 -14.114 61.335 109.749 1.00 2.00 0 ATOM 1871 RO RO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 63.396 113.089 1.00 2.00 0 ATOM 1876 C THR 193 -11.552 63.396 113.099 1.00 2.00 0 ATOM 1887 RO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1888 RO C THR 193 -11.552 63.396 113.099 1.00 2.00 0 ATOM 1888 RO C THR 193 -11.552 63.396 113.099 1.00 2.00 0 ATOM 1879 RO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1880 C THR 193 -11.552 63.396 113.099 1.00 2.00 0 ATOM 1879 RO 195 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1880 C THR 193 -11.552 63.396 113.099 1.00 2.00 0 ATOM 1880 C THR 193 -11.552 63.063 114.540 1.00 2.00 0 ATOM 1880 C THR 193 -11.552 63.063 114.540 1.00 2.00 0 ATOM 1880 C THR 193 -11.552 63.063 113.00 9.98 0 ATOM 1880 C PHD 195 -196 RO 196 RO 196 RO 196 RO 196 RO 196 RO 196 RO 19				-			60.194	115.015		
ATOM 1845 N ARG 191 -15.023 59.057 114.346 1.00 2.00 0 ATOM 1851 CA ARG 191 -13.732 58.738 113.772 1.00 1.00 2.00 0 ATOM 1852 CB ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1853 CG ARG 191 -12.300 56.900 112.821 1.00 11.31 0 ATOM 1854 CD ARG 191 -13.731 57.384 113.089 1.00 2.00 0 ATOM 1855 NE ARG 191 -13.139 54.935 113.762 1.00 11.31 0 ATOM 1857 CZ ARG 191 -13.139 54.935 113.762 1.00 12.00 0 ATOM 1858 NE ARG 191 -13.124 53.663 113.461 1.00 2.00 0 ATOM 1858 NENI ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1858 NENI ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1864 C ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1865 NE ARG 191 -13.084 59.769 112.857 1.00 2.00 0 ATOM 1866 N PRO 192 -13.685 60.267 113.202 1.00 6.10 0 ATOM 1866 N PRO 192 -13.685 60.065 111.070 1.00 37.06 0 ATOM 1866 CD PRO 192 -13.685 60.065 111.070 1.00 37.06 0 ATOM 1868 CA PRO 192 -13.685 60.065 111.070 1.00 37.06 0 ATOM 1869 CB PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1870 C PRO 192 -12.787 62.261 111.693 1.00 2.00 0 ATOM 1871 C PRO 192 -12.787 62.261 111.693 1.00 37.24 0 ATOM 1872 O PRO 192 -12.787 63.033 112.004 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 113.188 1.00 2.00 0 ATOM 1876 CB THR 193 -11.182 63.396 113.089 1.00 2.00 0 ATOM 1877 OGI THR 193 -11.627 63.063 113.194 1.00 2.00 0 ATOM 1887 OG THR 193 -11.627 63.063 113.194 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.194 1.00 2.00 0 ATOM 1881 O THR 193 -11.626 66.371 112.912 1.00 49.25 0 ATOM 1880 C ASF 194 -7.927 66.805 113.194 1.00 2.00 0 ATOM 1881 O THR 193 -10.862 61.859 115.089 1.00 2.00 0 ATOM 1889 C ASF 194 -7.927 66.805 113.194 1.00 49.25 0 ATOM 1889 C ASF 194 -7.927 66.805 113.194 1.00 49.25 0 ATOM 1889 C ASF 194 -7.927 66.818 110.049 1.00 2.00 0 ATOM 1889 C ASF 194 -7.927 66.818 110.049 1.00 2.00 0 ATOM 1889 C ASF 194 -7.927 66.818 110.049 1.00 2.00 0 ATOM 1889 C ASF 194 -7.927 66.818 110.049 1.00 2.00 0 ATOM 1889 C ASF 194 -7.927 66.818 110.049 1.00 2.00 0 ATOM 1899 C VAL 195 -5.971 65.948 64.611 116.839 1.00 2.00 0 ATOM 1899 C PRO 196 -6.874 66.15						-14.287	61.041	115.106	1.00 32.57	
ATOM 1851 CA ARG 191 -13.731 58.738 113.762 1.00 2.00 0 ATOM 1853 CG ARG 191 -12.300 56.900 112.821 1.00 11.31 0 ATOM 1854 CD ARG 191 -12.300 56.900 112.821 1.00 11.31 0 ATOM 1855 NE ARG 191 -13.1319 54.935 113.752 1.00 11.31 0 ATOM 1857 CZ ARG 191 -13.1319 54.935 113.752 1.00 11.31 0 ATOM 1858 NH1 ARG 191 -13.1319 54.935 113.762 1.00 2.00 0 ATOM 1858 NH1 ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1858 NH1 ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1866 N ARG 191 -11.995 53.085 113.047 1.00 2.00 0 ATOM 1866 N ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1866 N ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1866 N ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1866 N ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1866 N ARG 192 -14.914 69.99 112.857 1.00 2.00 0 ATOM 1866 CG PRO 192 -14.914 69.99 112.857 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.914 61.335 109.749 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1871 C PRO 192 -13.697 63.033 112.045 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1876 CB THR 193 -11.152 63.036 113.089 1.00 2.00 0 ATOM 1877 OGI THR 193 -11.152 63.036 113.089 1.00 2.00 0 ATOM 1877 OGI THR 193 -11.627 63.063 114.540 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 2.00 0 ATOM 1880 C THR 193 -10.862 61.859 115.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.865 65.041 113.572 1.00 2.00 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -7.686 65.041 115.216 1.00 2.00 0 ATOM 1880 C ASP 194 -7.686 65.041 115.216 1.00 2.00 0 ATOM 1880 C ASP 194 -7.686 65.041 115.216 1.00 2.00 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1890 O ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1890 C ASP 194 -7.686 65.041 115.216 1.00 2.00 2.00 0 ATOM 1890 C ASP 194 -7.686 66.181 119.							59.057	114.346		
ATOM 1853 CG ARG 191 -11.978 55.799 113.752 1.00 11.31 0 ATOM 1854 CD ARG 191 -11.978 55.799 113.752 1.00 2.00 0 ATOM 1855 NE ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1856 NI ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1858 NIH ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1861 NIE2 ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1865 O ARG 191 -11.985 97.69 112.857 1.00 2.00 0 ATOM 1866 C ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1866 N PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1866 N PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1866 C PRO 192 -13.084 61.068 110.799 1.00 36.28 0 ATOM 1869 CB PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.695 60.267 113.202 1.00 37.24 0 ATOM 1871 C PRO 192 -13.695 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1870 CG THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1880 C THR 193 -11.552 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.152 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.153 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -11.627 63.063 113.089 1.00 2.00 0 ATOM 1880 C THR 193 -10.665 62.755 112.860 1.00 17.62 0 ATOM 1880 C ASP 194 -6.566 67.977 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.566 67.197 112.91				-			58.738	113.772		
ATOM 1854 CD ARG 191 -11.978 55.799 113.752 1.00 11.31 0 ATOM 1855 NE ARG 191 -13.119 54.935 113.762 1.00 2.00 0 ATOM 1857 CZ ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1858 NH1 ARG 191 -11.955 53.085 113.0761 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -14.263 52.984 113.330 1.00 2.00 0 ATOM 1866 N PRO 191 -13.084 59.769 112.857 1.00 2.00 0 ATOM 1865 N PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1866 N PRO 192 -13.685 60.267 113.202 1.00 6.10 0 ATOM 1866 N PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1866 CA PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1868 CA PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1871 C PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1872 O PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1875 CA THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1876 CB THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1877 OG1 THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1879 CG2 THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C THR 193 -10.862 61.859 113.089 1.00 21.46 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1880 C ASP 194 -6.561 67.19				-			56.900	112.821		
ATOM 1857 CZ ARG 191 -13.139 54.935 113.462 1.00 2.00 0 ATOM 1857 CZ ARG 191 -13.124 53.663 113.416 1.00 2.00 0 ATOM 1858 NH1 ARG 191 -11.985 53.085 113.407 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1864 C ARG 191 -13.084 59.769 112.857 1.00 2.00 0 ATOM 1866 N PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1866 N PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1866 N PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1866 CD PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1869 CB PRO 192 -14.114 61.335 109.749 1.00 2.00 0 ATOM 1870 CG PRO 192 -12.787 62.261 111.693 1.00 2.00 0 ATOM 1871 C PRO 192 -12.787 62.261 111.693 1.00 2.00 0 ATOM 1872 O PRO 192 -13.697 63.033 112.004 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1875 CA THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1876 CB THR 193 -11.627 63.063 114.540 1.00 2.00 0 ATOM 1879 CG1 THR 193 -11.627 63.063 114.540 1.00 2.00 0 ATOM 1879 CG2 THR 193 -10.862 61.859 115.099 1.00 2.00 0 ATOM 1880 C TERR 193 -9.700 63.661 113.152 1.00 2.00 0 ATOM 1880 C TERR 193 -9.904 64.879 113.512 1.00 2.00 0 ATOM 1881 O THR 193 -9.394 64.889 115.099 1.00 2.00 0 ATOM 1888 OD ASP 194 -9.394 64.889 113.512 1.00 2.00 0 ATOM 1888 OD ASP 194 -6.551 67.197 112.912 1.00 49.25 0 ATOM 1888 OD ASP 194 -6.5920 66.371 112.217 1.00 49.25 0 ATOM 1889 C ASP 194 -6.5920 66.371 112.217 1.00 49.25 0 ATOM 1889 C CA PRO 196 -7.686 65.94 117.511 1.00 12.88 0 ATOM 1899 N VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1895 CG1 VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1895 CG2 VAL 195 -5.948 64.611 116.839 1.00 2.00 0 ATOM 1896 CG2 VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1897 CG1 VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1890 C PRO 196 -7.866 65.214 119.033 1.00 9.98 0 ATOM 1895 CG2 VAL 195 -5.948 64.611 116.839 1.00 9.98 0 ATOM 1895 CG1 VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1896 CG2 VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1895 CG1 VAL 195 -6.866 67.735 120.027 1.00 25.86 0 ATOM 1900 CD PRO 196 -6.862 67.							55.799	113.752		
ATOM 1857 CZ ARG 191 -11.985 53.085 113.047 1.00 2.00 0 ATOM 1861 NH2 ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1865 NH1 ARG 191 -14.263 52.984 113.430 1.00 2.00 0 ATOM 1865 O ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1865 O ARG 191 -11.995 60.267 113.202 1.00 6.10 0 ATOM 1866 N PRO 192 -13.685 60.065 111.670 1.00 37.06 0 ATOM 1866 CA PRO 192 -13.048 61.068 110.799 1.00 36.28 0 ATOM 1868 CA PRO 192 -14.912 59.530 111.045 1.00 2.00 0 ATOM 1868 CA PRO 192 -14.14 61.335 109.749 1.00 2.00 0 ATOM 1870 CG PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1871 C PRO 192 -14.743 59.973 109.602 1.00 2.00 0 ATOM 1873 N THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1875 CA THR 193 -11.552 62.348 112.168 1.00 2.00 0 ATOM 1876 CB THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1877 OGI THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1879 CG2 THR 193 -11.627 63.063 114.540 1.00 22.05 0 ATOM 1880 C THR 193 -10.862 61.859 115.089 1.00 22.05 0 ATOM 1880 C THR 193 -9.700 63.661 113.1572 1.00 21.46 0 ATOM 1881 O THR 193 -9.700 63.661 113.1572 1.00 21.00 0 ATOM 1882 N ASP 194 -9.394 64.879 113.572 1.00 21.00 0 ATOM 1886 CG ASP 194 -9.394 64.879 113.572 1.00 22.00 0 ATOM 1888 OD ASP 194 -9.394 64.879 113.572 1.00 20.00 0 ATOM 1888 OD ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1888 OD ASP 194 -7.927 66.805 113.431 1.00 49.25 0 ATOM 1888 OD ASP 194 -7.927 66.805 113.431 1.00 49.25 0 ATOM 1889 OD ASP 194 -6.561 67.197 112.912 1.00 49.25 0 ATOM 1889 OD ASP 194 -7.686 65.041 115.216 1.00 2.00 0 ATOM 1899 CG ASP 194 -7.927 66.805 113.431 1.00 49.25 0 ATOM 1899 OD ASP 194 -8.571 64.978 116.077 1.00 49.25 0 ATOM 1899 OD ASP 194 -7.5920 66.371 112.217 1.00 49.25 0 ATOM 1899 OD ASP 194 -8.571 64.978 116.077 1.00 49.25 0 ATOM 1899 OD ASP 194 -7.5920 66.371 112.217 1.00 49.25 0 ATOM 1899 OD ASP 194 -7.5920 66.371 112.217 1.00 49.25 0 ATOM 1899 OD ASP 194 -7.5920 66.371 112.217 1.00 49.25 0 ATOM 1899 OD ASP 194 -7.5920 66.371 112.217 1.00 49.25 0 ATOM 1899 OD ASP 194 -7.5920 66.371 112.217 1.00 49.25 0 ATOM 1899 OD ASP 194 -7			NE	ARG						
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ATOM 1880 C THR 193							61.859	115.089	1.00 21.46	
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ATOM 1889 C ASP 194 -7.686 65.041 115.216 1.00 54.62 0 ATOM 1890 O ASP 194 -8.571 64.978 116.077 1.00 54.62 0 ATOM 1891 N VAL 195 -6.408 64.836 115.488 1.00 17.01 0 ATOM 1893 CA VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1894 CB VAL 195 -4.509 64.017 116.818 1.00 9.98 0 ATOM 1895 CG1 VAL 195 -3.934 63.875 118.239 1.00 9.98 0 ATOM 1896 CG2 VAL 195 -4.532 62.675 116.124 1.00 9.98 0 ATOM 1897 C VAL 195 -5.971 65.994 117.511 1.00 12.88 0 ATOM 1898 O VAL 195 -5.343 66.934 117.038 1.00 9.98 0 ATOM 1899 N PRO 196 -6.771 66.152 118.567 1.00 16.08 0 ATOM 1900 CD PRO 196 -7.805 65.214 119.033 1.00 26.02 0 ATOM 1901 CA PRO 196 -6.862 67.429 119.284 1.00 21.91 0 ATOM 1902 CB PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1905 O PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1906 N ASP 197 -5.291 69.010 120.302 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0					-		66.371	112.217		
ATOM 1890 O ASP 194 -8.571 64.976 110.077 1.00 17.01 0 1891 N VAL 195 -6.408 64.836 115.488 1.00 17.01 0 12.88 0 1893 CA VAL 195 -5.948 64.611 116.839 1.00 12.88 0 1894 CB VAL 195 -4.509 64.017 116.818 1.00 9.98 0 1895 CG1 VAL 195 -4.509 64.017 116.818 1.00 9.98 0 1896 CG2 VAL 195 -3.934 63.875 118.239 1.00 9.98 0 1896 CG2 VAL 195 -5.971 65.994 117.511 1.00 12.88 0 1897 C VAL 195 -5.971 65.994 117.511 1.00 12.88 0 1897 C VAL 195 -5.343 66.934 117.038 1.00 9.98 0 1898 0 VAL 195 -5.343 66.934 117.038 1.00 9.98 0 1807 0 1908 CD PRO 196 -6.771 66.152 118.567 1.00 16.08 0 1908 CD PRO 196 -7.805 65.214 119.033 1.00 26.02 0 1807 0 1909 CD PRO 196 -6.862 67.429 119.284 1.00 21.91 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1904 C PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1904 C PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 23.54 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 23.54 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 23.54 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 23.54 0 1807 0 1908 CG PRO 196 -8.874 66.153 119.496 1.00 23.54 0 1807 0 18					194		65.041	115.216	1.00 2.00	
ATOM 1893 CA VAL 195 -5.948 64.611 116.839 1.00 12.88 0 ATOM 1894 CB VAL 195 -4.509 64.017 116.818 1.00 9.98 0 ATOM 1895 CG1 VAL 195 -3.934 63.875 118.239 1.00 9.98 0 ATOM 1896 CG2 VAL 195 -4.532 62.675 116.124 1.00 9.98 0 ATOM 1897 C VAL 195 -5.971 65.994 117.511 1.00 12.88 0 ATOM 1898 O VAL 195 -5.343 66.934 117.038 1.00 9.98 0 ATOM 1899 N PRO 196 -6.771 66.152 118.567 1.00 16.08 0 ATOM 1900 CD PRO 196 -7.805 65.214 119.033 1.00 26.02 0 ATOM 1901 CA PRO 196 -6.862 67.429 119.284 1.00 21.91 0 ATOM 1902 CB PRO 196 -6.862 67.195 120.240 1.00 25.86 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1905 O PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1906 N ASP 197 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0			0				64.978 64.836	115.488	1.00 17.01	
ATOM 1894 CB VAL 195							64.611	116.839	1.00 12.88	
ATOM 1895 CG1 VAL 195 -3.934 63.875 118.239 1.00 9.98 0 ATOM 1896 CG2 VAL 195 -4.532 62.675 116.124 1.00 9.98 0 ATOM 1897 C VAL 195 -5.971 65.994 117.511 1.00 12.88 0 ATOM 1898 O VAL 195 -5.343 66.934 117.038 1.00 9.98 0 ATOM 1899 N PRO 196 -6.771 66.152 118.567 1.00 16.08 0 ATOM 1900 CD PRO 196 -7.805 65.214 119.033 1.00 26.02 0 ATOM 1901 CA PRO 196 -6.862 67.429 119.284 1.00 21.91 0 ATOM 1902 CB PRO 196 -8.026 67.195 120.240 1.00 25.86 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1905 O PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1906 N ASP 197 -4.809 66.818 120.349 1.00 23.54 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 39.93 0						-4.509		116.818	1.00 9.98	
ATOM 1896 CG2 VAL 195 ATOM 1897 C VAL 195 ATOM 1898 O VAL 195 ATOM 1899 N PRO 196 ATOM 1900 CD PRO 196 ATOM 1901 CA PRO 196 ATOM 1902 CB PRO 196 ATOM 1903 CG PRO 196 ATOM 1903 CG PRO 196 ATOM 1904 C PRO 196 ATOM 1905 O PRO 196 ATOM 1905 O PRO 196 ATOM 1905 CA ASP 197 ATOM 1908 CA ASP 197	MOTA	1895								
ATOM 1898 O VAL 195 -5.343 66.934 117.038 1.00 9.98 0 ATOM 1899 N PRO 196 -6.771 66.152 118.567 1.00 16.08 0 ATOM 1900 CD PRO 196 -7.805 65.214 119.033 1.00 26.02 0 ATOM 1901 CA PRO 196 -6.862 67.429 119.284 1.00 21.91 0 ATOM 1902 CB PRO 196 -8.026 67.195 120.240 1.00 25.86 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1905 O PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1905 O PRO 196 -4.809 66.818 120.349 1.00 23.54 0 ATOM 1906 N ASP 197 -4.809 66.818 120.349 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0										0
ATOM 1899 N PRO 196 -6.771 66.152 118.567 1.00 16.08 0 ATOM 1900 CD PRO 196 -7.805 65.214 119.033 1.00 26.02 0 ATOM 1901 CA PRO 196 -6.862 67.429 119.284 1.00 21.91 0 ATOM 1902 CB PRO 196 -8.026 67.195 120.240 1.00 25.86 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1905 O PRO 196 -4.809 66.818 120.349 1.00 23.54 0 ATOM 1906 N ASP 197 -5.291 69.010 120.302 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0								117.038		
ATOM 1900 CD PRO 196 -7.805 65.214 119.284 1.00 21.91 0 ATOM 1901 CA PRO 196 -6.862 67.429 119.284 1.00 25.86 0 ATOM 1902 CB PRO 196 -8.026 67.195 120.240 1.00 25.86 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1905 O PRO 196 -4.809 66.818 120.349 1.00 23.54 0 ATOM 1906 N ASP 197 -4.809 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0					196					
ATOM 1901 CA PRO 196 -8.026 67.195 120.240 1.00 25.86 0 ATOM 1902 CB PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1905 O PRO 196 -4.809 66.818 120.349 1.00 23.54 0 ATOM 1906 N ASP 197 -5.291 69.010 120.302 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0	ATOM	1900								
ATOM 1903 CG PRO 196 -8.874 66.153 119.496 1.00 25.31 0 ATOM 1904 C PRO 196 -5.560 67.735 120.027 1.00 24.64 0 ATOM 1905 O PRO 196 -4.809 66.818 120.349 1.00 23.54 0 ATOM 1906 N ASP 197 -5.291 69.010 120.302 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0									1.00 25.86	0
ATOM 1904 C PRO 196 -5.560 67.735 120.027 1.00 24.64 O ATOM 1905 O PRO 196 -4.809 66.818 120.349 1.00 23.54 O ATOM 1906 N ASP 197 -5.291 69.010 120.302 1.00 39.93 O ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 O ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 O ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 78.93 O						-8.874	66.153	119.496		
ATOM 1905 O PRO 190 ATOM 1906 N ASP 197 -5.291 69.010 120.302 1.00 39.93 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0 ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 78.93 0				PRO	196					
ATOM 1908 CA ASP 197 -4.058 69.377 121.005 1.00 38.76 0									1.00 39.93	
AION 1900 CA 101 107 3 605 70 789 120 612 1 00 78 93 0							69.377	121.005	1.00 38.76	
							70.789	120.612	1.00 78.93	O

				107	-4.726	71 900	120.658	1.00 84.50	0
MOTA MOTA	1910 1911	CG A	SP	197 197	-5.071	72.263	121.768	1.00 83.41	ŏ
ATOM	1912		SP	197	-5.258	72.134	119.578	1.00 90.03	0
ATOM	1913	_	SP	197	-4.162		122.520	1.00 41.84	0
MOTA	1914		SP	197	-3.227 -5.316		123.255 122.973	1.00 85.35 1.00 2.00	0
MOTA	1915 1917		LN LN	198 198	-5.581	68.588	124.385	1.00 2.00	ŏ
MOTA MOTA	1918		LN	198	-5.686	69.937	125.120	1.00 27.32	Ō
MOTA	1919		LN	198	-6.621	70.967	124.503	1.00 32.39	0
MOTA	1920		LN	198	-7.981		125.192	1.00 29.94	0
MOTA	1921		LN	198	-8.108 -9.007		126.397 124.424	1.00 32.81 1.00 29.42	0
ATOM	1922	_	iln Ln	198 198	-6.863		124.492	1.00 2.00	ŏ
MOTA MOTA	1925 1926	_	LN	198	-7.748	67.909	123.640	1.00 26.90	ŏ
ATOM	1927		LY	199	-6.943	66.949	125.509	1.00 15.20	0
MOTA	1929	CA G	LY	199	-8.137	66.148	125.692	1.00 15.56	Ŏ
MOTA	1930		LY	199	-7.820		125.631 125.502	1.00 13.32 1.00 2.00	0
MOTA	1931		LY	199	-6.655 -8.857		125.684	1.00 2.00	ő
MOTA	1932		'en 'en	200 200	-8.690	62.402	125.666	1.00 2.00	Õ
MOTA MOTA	1934 1935		EU.	200	-10.049	61.701	125.667	1.00 2.00	0
ATOM	1936		EU	200	-9.945	60.224	126.081	1.00 2.00	0
ATOM	1937	CD1 L		200	-11.126		126.947	1.00 2.00	0
MOTA	1938	CD2 L		200	-9.842		124.847 124.521	1.00 2.00 1.00 2.00	0
ATOM	1939	_	EU	200 200	-7.850 -6.764		124.764	1.00 2.00	ŏ
ATOM ATOM	1940 1941	-	.EU .EU	201	-8.344	62.045	123.283	1.00 2.00	0
ATOM	1943	-	EU	201	-7.663	61.566	122.070	1.00 2.00	0
ATOM	1944		EU	201	-8.378		120.811	1.00 2.00	0
MOTA	1945		EU	201	-B.369		119.589 118.375	1.00 2.00 1.00 2.00	0
ATOM	1946	CD1 L		201	-8.895 -6.981	60 569	119.277	1.00 2.00	ŏ
ATOM	1947 1948	CD2 L	'Ea 'Ea	201 201	-6.246	62.078	122.059	1.00 2.00	0
ATOM ATOM	1949	-	EU	201	-5.319	61.347	121.715	1.00 2.00	0
ATOM	1950		YS	202	-6.071	63.333		1.00 2.00	0
MOTA	1952		:YS	202	-4.733		122.484 123.024	1.00 2.00 1.00 2.00	0
MOTA	1953		TYS	202 202	-4.722 -3.012		123.024	1.00 2.00	Õ
ATOM ATOM	1954 1955		YS YS	202	-3.850		123.394	1.00 2.00	0
ATOM	1956		YS	202	-2.719	62.715	123.036	1.00 2.00	0
ATOM	1957		SP	203	-4.385		124.584	1.00 15.07 1.00 11.86	0
MOTA	1959	_	ASP	203	-3.673	62.057 62.142	125.611 126.931	1.00 11.86 1.00 13.71	ő
MOTA	1960		ASP	203 203	-4.429 -4.449		127.497	1.00 15.77	0
MOTA MOTA	1961 1962	CG A	ASP ASP	203	-3.529	64.315	127.162	1.00 11.73	0
ATOM	1963	OD2 A		203	-5.380	63.861	128.271	1.00 16.16	0
MOTA	1964	C A	ASP	203	-3.369	60.615	125.262	1.00 13.81 1.00 23.94	0
MOTA	1965		ASP	203	-2.254	60.155 59.902	125.492 124.695	1.00 10.49	Ö
MOTA	1966		LBU LBU	204 204	-4.330 -4.097	58.502	124.332	1.00 10.49	0
MOTA MOTA	1968 1969		LEU	204	-5.351	57.912		1.00 10.22	0
MOTA	1970		LEU	204	-6.47B			1.00 10.22	0
MOTA	1971	CD1 I		204	-7.667	57.281	123.866 125.620	1.00 10.22 1.00 10.22	Ö
MOTA	1972	CD2 I		204	-6.143 -2.930	56.441 58.347	123.361	1.00 10.49	Ö
MOTA	1973 1974		Leu Leu	204 204	-2.204	57.352	123.392	1.00 10.22	0
MOTA MOTA	1975		LEU	205	-2.748	59.336	122.501	1.00 2.00	0
ATOM	1977		LEU	205	-1.679	59.301	121.525	1.00 2.00	0
MOTA	1978	CB I	LEU	205	-2.161	59.933	120.219 119.551	1.00 2.00 1.00 2.00	0
ATOM	1979		LEU	205	-3.459 -3.595	59.463 60.238	119.331	1.00 2.00	Õ
ATOM	1980 1981	CD1 I		205 205	-3.459	57.954	119.269	1.00 2.00	0
MOTA MOTA	1982		LEU	205	-0.409	60.018		1.00 2.00	0
ATOM	1983		LEU	205	0.620	59.890		1.00 2.00	0
ATOM	1984	_	TRP	206	-0.472	60.756 61.526		1.00 10.02 1.00 10.02	0
MOTA	1986	_	TRP	206 206	0.678 0.362	62.994		1.00 2.00	ŏ
MOTA MOTA	1987 1988		TRP TRP	206	0.413	63.485		1.00 2.00	0
A LON	¥ 7 U U								

					c4 102 121 420	1.00 2.00	0
MOTA	1989	CD2 TRP	206	1.532 1.105	64.103 121.420 64.535 120.148	1.00 2.00	Ö
MOTA	1990	CE2 TRP	206 206	2.854	64.355 121.813	1.00 2.00	0
MOTA MOTA	1991 1992	CD1 TRP	206	-0.616	63.541 121.162	1.00 2.00	0
ATOM	1993	NE1 TRP	206	-0.212	64.178 120.017	1.00 2.00	0
ATOM	1995	CZ2 TRP	206	1.950	65.207 119.263	1.00 2.00 1.00 2.00	0
MOTA	1996	CZ3 TRP	206	3.697	65.024 120.932	1.00 2.00 1.00 2.00	0
MOTA	1997	CH2 TRP	206	3.238	65.443 119.669 61.330 124.955	1.00 10.02	ŏ
MOTA	1998	C TRP	206	1.246 2.419	61.639 125.194	1.00 2.00	ŏ
MOTA	1999	O TRP	206 207	0.418	60.867 125.886	1.00 2.00	0
MOTA	2000	n ser Ca ser	207	0.830	60.690 127.267	1.00 2.00	0
MOTA	2002 2003	CA SER CB SER	207	-0.363	60.273 128.121	1.00 29.76	0
MOTA MOTA	2004	OG SER	207	-0.482	61.120 129.256	1.00 35.73	0
ATOM	2006	C SER	207	1.951	59.699 127.446	1.00 2.00	0
MOTA	2007	O SER	207	2.075	58.743 126.681	1.00 31.05 1.00 2.00	Ö
MOTA	2008	n asp	208	2.777	59.943 128.462 59.056 128.764	1.00 2.00	Ö
MOTA	2010	CA ASP	208	3.899 5.2 5 7	59.714 128.505	1.00 42.39	ŏ
MOTA	2011	CB ASP	208 208	5.296	60.532 127.256	1.00 46.59	0
MOTA	2012	CG ASP OD1 ASP	208	5.695	61.706 127.362	1.00 48.93	0
MOTA	2013 2014	OD2 ASP	208	4.959	60.008 126.180	1.00 44.70	0
MOTA MOTA	2015	C ASP	208	3.903	58.705 130.231	1.00 2.00	0
ATOM	2016	O ASP	208	3.580	59.537 131.072	1.00 45.54	0
ATOM	2017	N PRO	209	4.244	57.457 130.559	1.00 10.98 1.00 2.00	0
ATOM	2018	CD PRO	209	4.509	56.316 129.673 57.040 131.957	1.00 10.98	ŏ
MOTA	2019	CA PRO	209	4.299 4.407	55.522 131.845	1.00 2.00	ŏ
MOTA	2020	CB PRO	209	5.197	55.356 130.604	1.00 2.00	O
MOTA	2021	CG PRO	209 209	5.584	57.675 132.507	1.00 10.98	0
MOTA MOTA	2022 2023	C PRO O PRO	209	6.448	58.078 131.712	1.00 2.00	0
ATOM	2024	N ASP	210	5.724	57.774 133.833	1.00 2.00	0
ATOM	2026	CA ASP	210	6.931	58.360 134.413	1.00 2.00	0
ATOM	2027	CB ASP	210	6.755	59.863 134.614	1.00 31.00	Ö
ATOM	2028	CG ASP	210	8.050	60.570 134.990 59.956 134.874	1.00 33.32	Ö
MOTA	2029	OD1 ASP	210	9.132 7.989	61.756 135.396	1.00 38.93	Ō
MOTA	2030	OD2 ASP	210	7.304	57.734 135.729	1.00 2.00	0
MOTA	2031	C ASP O ASP	210 210	6.448	57.320 136.496	1.00 28.60	0
MOTA MOTA	2032 2033	O ASP N LYS	211	8.608		1.00 4.23	0
ATOM	2035	CA LYS	211	9.170	57.136 137.211	1.00 9.37	0
MOTA	2036	CB LYS	211	10.647	56.760 137.024	1.00 17.66 1.00 29.60	Ö
MOTA	2037	CG LYS	211	10.941	55.441 136.345 55.184 136.403	1.00 32.82	ŏ
MOTA	2038	CD LYS	211	12.451 12.852	55.184 136.403 53.831 135.809	1.00 42.70	ō
MOTA	2039	CE LYS	211 211	14.336	53.616 135.852	1.00 41.45	0
ATOM	2040	NZ LYS	211	9.100	58.157 138.367	1.00 7.69	0
MOTA MOTA	2044 2045	C LYS	211	8.746	57.803 139.481	1.00 18.85	Ō
ATOM	2046	N ASP	212	9.456	59.414 138.093	1.00 2.00	0
MOTA	2048	CA ASP	212	9.468	60.483 139.101	1.00 2.00 1.00 39.78	ő
MOTA	2049	CB ASP	212	10.322	61.669 138.624 61.254 137.754	1.00 46.31	Ö
MOTA	2050	CG ASP	212	11.483 11.823	62.030 136.834	1.00 41.85	0
MOTA	2051	OD1 ASP OD2 ASP	212 212	12.051	60.166 137.987	1.00 46.28	0
MOTA	2052 2 053	OD2 ASP		8.074	61.033 139.412	1.00 2.00	0
MOTA MOTA	2054	0 ASP	212	7.943	62.164 139.903	1.00 41.82	0
MOTA	2055	N VAL	213	7.040	60.250 139.124	1.00 2.00	0
MOTA	2057	CA VAL	213	5.669	60.690 139.340	1.00 2.00 1.00 2.00	0
ATOM	205B	CB VAL	213	5.137	61.376 138.050 61.247 137.922	1.00 2.00	ŏ
ATOM	2059	CG1 VAL	213	3.652	61.247 137.922 62.826 138.076	1.00 2.00	Ŏ
ATOM	2060	CG2 VAL	213	5.505 4.767	59.526 139.763	1.00 2.00	0
MOTA	2061	C VAL	213 213	5.007	58.356 139.390	1.00 2.00	0
ATOM	2062	O VAL N LEU	213	3.748	59.844 140.564	1.00 21.53	0
MOTA	2063 2065	N LEU CA LEU	214	2.815	58.830 141.022	1.00 19.34	0
MOTA MOTA	2066	CB LEU	214	2.742	58.809 142.543	1.00 14.66	0
ATOM	2067	CG LEU	214	2.371	57.438 143.097	1.00 15.86 1.00 9.71	0
ATOM	2068	CD1 LEU	214	3.507	56.477 142.816	1.00 9.71	U

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MOTA MOTA	2069 2070	CD2 C	LEU LEU	214 214	2.109 1.433		144.588 140.439	1.00 17.44 1.00 19.87	0
ATOM	2071	ŏ	LEU	214	0.757		140.033	1.00 9.54	0
MOTA	2072	N	GLY	215 215	1.003 -0.299	60.322 60.640	140.411 139.852	1.00 16.98 1.00 11.19	0
MOTA MOTA	2074 2075	CA C	GLY GLY	215	-0.081		138.487	1.00 12.07	ō
ATOM	2076	ō	GLY	215	0.772		137.739	1.00 10.02	0
MOTA	2077	N	TRP	216	-0. 824 -0.684		138.167 136.890	1.00 2.00 1.00 2.00	0
ATOM ATOM	2079 2080	CA CB	TRP TRP	216 216	-2.030		136.458	1.00 2.00	Õ
ATOM	2081	CG	TRP	216	-2.903	62.452	136.050	1.00 2.00	0
MOTA	2082	CD2	TRP	216	-2.840		134.804	1.00 2.00 1.00 2.00	0
MOTA	2083 2084	CE2	TRP TRP	216 216	-3.806 -2.050		134.865 133.648	1.00 2.00	ŏ
MOTA MOTA	2085	CDI	TRP	216	-3.879	61.860	136.789	1.00 2.00	0
ATOM	2086	NE1	TRP	216	-4.425		136.087	1.00 2.00	0
MOTA	2088	CZ2	TRP	216	-4.005 -2.251		133.809 132.605	1.00 2.00 1.00 2.00	0
MOTA	2089 2090	CZ3 CH2	TRP TRP	216 216	-3.217		132.695	1.00 2.00	ŏ
ATOM ATOM	2091	C	TRP	216	0.310	64.101	136.985	1.00 2.00	0
MOTA	2092	0	TRP	216	0.261		137.926	1.00 2.00 1.00 12.63	0
MOTA	2093	N	GLY GLY	217 217	1.213 2.199		136.021 136.028	1.00 15.52	ŏ
ATOM ATOM	2095 2096	CA C	GLY	217	2.312	65.928	134.675	1.00 10.15	0
ATOM	2097	ō	GLY	217	1.627		133.720	1.00 26.84 1.00 23.86	0
ATOM	2098	N	GLU	218	3.189 3.434		134.594 133.364	1.00 23.80	ŏ
MOTA	2100 2101	CA CB	GLU	218 218	4.198	68.978	133.729	1.00 65.82	0
MOTA MOTA	2101	œ	GLU	218	5.082	69.59 5	132.641	1.00 67.96	0
MOTA	2103	CD	GLU	218	6.577		132.987 133.631	1.00 67.00 1.00 68.41	0
MOTA	2104	OE1	GLU GLU	218 218	7.056 7.2 7 6	68.589	132.616	1.00 63.29	ŏ
MOTA MOTA	2105 2106	OE2 C	GLU	218	4.232	66.841	132.371	1.00 25.48	0
ATOM	2107	ŏ	GLU	218	4.808		132.750	1.00 67.07 1.00 21.19	0
ATOM	2108	N	ASN	219 219	4.253 5.039		131.103 130.108	1.00 21.13	ŏ
MOTA MOTA	2110 2111	CA CB	asn Asn	219	4.140	65.888	129.048	1.00 10.60	0
MOTA	2112	ČĞ	ASN	219	4.832	64 701	120 202		
MOTA							128.282	1.00 14.78	0
	2113	OD1	ASN	219	6.052	64.646	128.341	1.00 14.78 1.00 9.20 1.00 8.28	0
MOTA	2114	OD1 ND2	asn Asn	219 219		64.646 64.003 67.411	128.341 127.560 129.425	1.00 9.20 1.00 8.28 1.00 21.70	0 0 0
		OD1	ASN	219 219 219 219 219	6.052 4.057 6.058 5.776	64.646 64.003 67.411 68.576	128.341 127.560 129.425 129.151	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52	0 0 0
ATOM ATOM ATOM ATOM	2114 2117 2118 2119	OD1 ND2 C O N	asn asn asn asn asp	219 219 219 219 219 220	6.052 4.057 6.058 5.776 7.246	64.646 64.003 67.411 68.576 66.886	128.341 127.560 129.425 129.151 129.151	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32	0 0 0
ATOM MOTA ATOM ATOM MOTA	2114 2117 2118 2119 2121	OD1 ND2 C O N CA	ASN ASN ASN ASN ASP ASP	219 219 219 219 219 220 220	6.052 4.057 6.058 5.776 7.246 8.266	64.646 64.003 67.411 68.576 66.886 67.690	128.341 127.560 129.425 129.151	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55	0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122	OD1 ND2 C O N CA CB	asn asn asn asn asp	219 219 219 219 219 220	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673	128.341 127.560 129.425 129.151 129.151 128.485 128.358 127.447	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38	0 0 0 0 0 0
ATOM MOTA ATOM ATOM MOTA	2114 2117 2118 2119 2121 2122 2123 2124	OD1 ND2 C O N CA CB CG OD1	ASN ASN ASN ASP ASP ASP ASP ASP	219 219 219 219 220 220 220 220 220	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.083	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584	128.341 127.560 129.425 129.151 129.151 128.485 128.358 127.447 127.938	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43	0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125	OD1 ND2 C O N CA CB CG OD1 OD2	ASN ASN ASN ASP ASP ASP ASP ASP	219 219 219 219 220 220 220 220 220 220	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.083 9.778	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148	128.341 127.560 129.425 129.151 129.151 128.485 128.358 127.447 127.938 126.237 127.100	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38	00000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126	OD1 ND2 C O N CA CB CG OD1 OD2 C	ASN ASN ASN ASP ASP ASP ASP ASP	219 219 219 219 220 220 220 220 220	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.083	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262	128.341 127.560 129.425 129.151 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 34.19 1.00 48.55 1.00 92.38 1.00 92.38 1.00 92.09 1.00 33.44 1.00 47.84	00000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128	OD1 ND2 C O N CA CB CG OD1 OD2	ASN ASN ASN ASP ASP ASP ASP ASP ASP ASP ASP	219 219 219 219 220 220 220 220 220 220 220 220 220 22	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.778 7.763 B.045 6.986	64.646 64.003 67.411 68.576 66.886 67.690 65.673 64.584 65.797 68.148 69.262 67.287	128.341 127.560 129.425 129.151 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.444	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130	OD1 ND2 C O N CA CB CG OD1 OD2 C	ASN ASN ASN ASP ASP ASP ASP ASP ASP ASP ASP ASP ASP	219 219 219 219 220 220 220 220 220 220 220 221 221	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.083 9.778 7.763 8.045 6.986	64.646 64.003 67.411 68.576 66.886 67.690 65.673 64.584 65.797 68.148 69.262 67.287 67.552	128.341 127.560 129.425 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.444 125.127	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 34.19 1.00 48.55 1.00 92.38 1.00 92.38 1.00 92.09 1.00 33.44 1.00 47.84	00000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131	OD1 ND2 C O N CA CB OD1 OD2 C O N CA CB	ASN ASN ASN ASP ASP ASP ASP ASP ASP ASP ASP ASP ASP	219 219 219 219 220 220 220 220 220 220 220 220 220 22	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.778 7.763 B.045 6.986	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262 67.287 67.552 66.382 65.056	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98 1.00 2.11 1.00 2.00 1.00 2.00	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130	OD1 ND2 C O N CA CB CG OD1 OD2 C	ASN ASN ASN ASP ASP ASP ASP ASP ASP ASP ASP ASP ASP	219 219 219 220 220 220 220 220 220 221 221 221 221	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.083 9.778 7.763 8.045 6.945 6.927 5.251	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262 67.287 67.552 66.382 65.056 63.981	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739 124.421	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 2.98 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2132 2133 2134	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CG OD1	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 220 221 221 221 221	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.778 7.763 8.045 6.986 6.427 5.578 5.241	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262 67.287 67.552 66.382 65.056 63.981 62.658	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739 124.421 124.590	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98 1.00 2.11 1.00 2.00 1.00 2.00	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2132 2133 2134 2136	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CCD N CA CB CCD N CA CB CCD N CA CB CCD N CA CCD CCD N N CCD N N N CCD N N CCD N N N N	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 220 221 221 221 221	6.052 4.057 6.058 5.776 7.246 8.266 9.585 9.469 9.083 9.778 8.045 6.986 6.427 5.251 5.241 5.2828 6.785	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262 67.287 67.552 66.382 65.056 63.981	128.341 127.560 129.425 129.151 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739 124.421 124.590 123.811 122.790	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2132 2133 2134 2136 2137	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CG OD1	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 220 221 221 221 221	6.052 4.057 6.058 5.7746 8.266 9.585 9.469 9.778 7.763 8.986 6.427 5.578 6.251 5.281 5.285	64.646 64.003 67.411 68.576 66.886 67.690 65.673 64.584 65.797 68.148 69.262 67.287 67.287 66.382 65.056 63.981 62.658 62.154 60.947	128.341 127.560 129.425 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.661 126.444 125.127 124.672 124.739 124.421 124.590 123.811 122.790 124.064	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98 1.00 22.11 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2132 2133 2134 2136 2137 2140 2143	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CC N CB CG N CA CB CG N CB CG N CB CG N CB CG N CB CG N CB CD N CB CB CD N CB CB CB CB CB CB CB CB CB CB CB CB CB	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.058 5.776 8.266 9.585 9.469 9.083 9.778 7.763 8.045 6.427 5.578 6.221 5.828 7.285 5.241	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262 67.287 67.552 66.382 65.056 63.981 62.858 62.154 62.8658 62.8658	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 125.127 124.672 124.739 124.421 124.590 123.811 122.790 124.064 125.109	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98 1.00 22.11 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2130 2131 2132 2133 2134 2136 2137 2140 2143 2144	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CC N CB CC N CB CC N CB CC N CB CC N CB CC N CB CC N CD N CD	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.058 5.776 8.246 9.585 9.469 9.083 9.778 7.763 8.045 6.427 5.578 5.241 5.828 6.785 7.255 5.164	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.673 64.584 65.797 68.148 69.262 67.287 67.552 66.356 63.981 62.658 62.154 62.866 60.947 69.239	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739 124.421 124.590 123.811 122.790 124.064 125.109 124.052	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2130 2131 2132 2133 2134 2136 2137 2140 2143 2144 2145	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CC NECZ NH1 NH2 C O N	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.058 5.776 8.266 9.469 9.083 9.7763 8.045 6.927 5.241 5.241 5.2546 5.2546 5.2546 5.188 4.328	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.797 68.148 69.287 67.552 67.287 67.552 63.981 62.658 62.658 62.658 62.658 62.658 63.981 69.239 69.239 69.247	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739 124.421 124.590 123.811 122.790 124.064 125.109 124.080 124.080 126.343	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2130 2131 2132 2133 2134 2136 2137 2140 2143 2144	OD1 ND2 C O N CA CB CG OD1 OD2 C O N CA CB CC N CB CC N CB CC N CB CC N CB CC N CB CC N CB CC N CD N CD	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.058 5.776 8.266 9.585 9.083 9.7763 8.045 6.251 5.241 5.241 5.251 5.241 5.285 7.285 5.188 4.328 2.939	64.646 64.003 67.411 68.576 66.886 67.690 66.900 65.573 65.797 68.148 69.267 67.287 67.552 66.382 65.0581 62.866 62.866 60.947 69.239 69.247 70.443	128.341 127.560 129.425 129.151 129.151 128.485 127.447 127.938 126.237 127.100 126.661 126.444 125.127 124.672 124.739 124.739 124.064 125.109 124.064 125.109 124.052 126.343 125.844	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 47.84 1.00 22.98 1.00 22.11 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2133 2134 2136 2137 2140 2143 2144 2145 2144 2144 2144 2147	OD1 ND2 C O N C C C C C C C C C C C C C C C C C	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.0558 5.7746 89.585 9.469 9.763 86.987 7.0486 6.427 5.578 5.251 5.8285 7.2285 5.241 86.927 5.164 5.188 4.328 2.142	64.646 64.003 67.411 68.576 66.886 67.690 65.673 64.584 65.797 68.148 69.262 67.252 67.252 65.056 63.981 62.856 63.981 62.858 62.858 63.981 62.858 63.981 69.280 70.447 70.198	128.341 127.560 129.425 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.661 126.444 125.127 124.672 124.739 124.421 124.590 123.811 125.109 124.064 125.109 124.052 126.280 126.344 125.509	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2132 2133 2134 2136 2140 2143 2144 2145 2145 2147 2148 2149 2150	OD1 ND2 C O N A CB CCD OD2 C O N A CB CCD NE C NH12 C O N C O N C O N C O N	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG GLY GLY VAL	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.058 5.776 8.266 9.585 9.469 9.083 9.778 7.763 8.045 6.427 5.578 6.2241 5.828 6.785 7.285 5.164 5.188 4.328 2.939 2.1458 1.356	64.646 64.003 67.411 68.576 66.886 67.690 65.673 64.584 65.797 68.148 69.262 67.287 67	128.341 127.560 129.425 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.661 124.672 124.739 124.672 124.739 124.672 124.739 124.064 125.127 124.052 126.280 126.343 125.809 125.776 125.342	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 33.44 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2114 2117 2118 2119 2121 2122 2123 2124 2125 2126 2127 2128 2130 2131 2133 2134 2136 2137 2140 2143 2144 2145 2144 2144 2144 2147	OD1 ND2 C O N C C C C C C C C C C C C C C C C C	ASN ASN ASN ASP ASP ASP ASP ASP ASP ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	219 219 219 220 220 220 220 220 221 221 221 221 221	6.052 4.057 6.058 5.776 8.2466 9.585 9.469 9.083 9.778 7.763 6.427 5.578 6.421 5.578 6.221 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.241 5.246 5.241 5.241 5.246 5.241 5.246 5.241 5.246 5.241 5.246 5.241 5.241 5.246 5.241 5.246 5.241 5.241 5.246 5.241 5.246 5.241 5.246 5.241 5.246 5.241 5.246 5.246 5.241 5.246 5.241 5.246 5.241 5.246 5.246 5.241 5.246	64.646 64.003 67.411 68.576 66.886 67.690 65.673 64.584 65.797 68.148 69.262 67.552 66.382 65.056 63.981 62.658 62.156 63.981 62.658 62.158 62.158 69.289 69	128.341 127.560 129.425 129.151 128.485 128.358 127.447 127.938 126.237 127.100 126.661 126.661 124.672 124.739 124.672 124.739 124.672 124.739 124.064 125.127 124.052 126.280 126.343 125.809 125.776 125.342	1.00 9.20 1.00 8.28 1.00 21.70 1.00 9.52 1.00 35.32 1.00 34.19 1.00 48.55 1.00 92.38 1.00 48.43 1.00 92.09 1.00 2.00	000000000000000000000000000000000000000

MOTA MOTA	2154 2155	CG1 VAL	223 223 223	-0.063 2.258 1.071	67.888 123.280 68.762 123.004 67.125 126.250	1.00 2.00 1.00 2.00 1.00 2.00	0 0 0
MOTA	2156	O VAL	223 223	2.004	66.431 126.656	1.00 2.00	0
MOTA	2157 2158	O VAL N SER	224	-0.205	66.897 126.550	1.00 12.21	0
MOTA MOTA	2160	CA SER	224	-0.661	65.825 127.440	1.00 15.55	0
ATOM	2161	CB SER	224	-0.609	64.453 126.759	1.00 6.64	0
ATOM	2162	OG SER	224	0.713	64.003 126.546 65.779 128.787	1.00 10.80 1.00 20.53	0
MOTA	2164	C SER	224	0.077	66.706 129.153	1.00 20.33	Õ
MOTA	2165	O SER	224 225	0.814 -0.124	64.700 129.533	1.00 6.68	Ö
MOTA	2166	N PHE	225	0.484	64.586 130.839	1.00 6.68	0
MOTA	2168 2169	CA PHE	225	-0.625	64.489 131.890	1.00 14.03	0
MOTA MOTA	2170	CG PHE	225	-1.670	65.552 131.758	1.00 11.21	0
ATOM	2171	CD1 PHE	225	-2.925	65.248 131.239	1.00 11.66	0
ATOM	2172	CD2 PHE	225	-1.396	66.869 132.131	1.00 11.41 1.00 11.76	0
ATOM	2173	CE1 PHE	225	-3.899 -2.363	66.241 131.086 67.869 131.983	1.00 14.72	ŏ
ATOM	2174	CE2 PHE	225 225	-3.618	67.553 131.458	1.00 16.73	ō
MOTA	2175	CZ PHE	225	1.405	63.392 130.957	1.00 6.68	0
MOTA MOTA	2176 2177	O PHE	225	1.902	62.850 129.966	1.00 15.31	0
ATOM	2178	N THR	226	1.663	63.022 132.203	1.00 2.00	0
ATOM	2180	CA THR	226	2.473	61.872 132.509	1.00 2.00	0
ATOM	2181	CB THR	226	3.898	62.226 132.853	1.00 13.51 1.00 13.51	0
ATOM	2182	OG1 THR	226	4.607	61.012 133.094 63.081 134.086	1.00 13.51	ő
MOTA	2184	CG2 THR	226	3.961 1.809	61.217 133.686	1.00 2.00	ŏ
ATOM	2185 2186	C THR O THR	226 226	1.039	61.861 134.394	1.00 13.51	0
MOTA MOTA	2187	N PHE	227	2.086	59.938 133.883	1.00 2.00	0
ATOM	2189	CA PHE	227	1.466	59.191 134.957	1.00 2.00	0
MOTA	2190	CB PHE	227	0.206	58.448 134.472 57.793 133.131	1.00 18.74 1.00 21.51	Ö
MOTA	2191	CG PHE	227	0.362 0.717	56.453 133.034	1.00 18.55	Ö
ATOM	2192	CD1 PHE	227 227	0.188	58.532 131.954	1.00 20.45	Ō
MOTA MOTA	2193 2194	CD2 PHE	227	0.902	55.857 131.787	1.00 19.97	0
ATOM	2195	CE2 PHE	227	0.371	57.948 130.698	1.00 14.44	0
MOTA	2196	CZ PHE	227	0.729	56.612 130.612	1.00 17.09 1.00 2.00	0
MOTA	2197	C PHE	227	2.427 3.282	58.224 135.562 57.668 134.888	1.00 20.11	Ö
MOTA	2198	O PHE	227 228	2.294		1.00 21.90	0
MOTA MOTA	2199 2201	N GLY CA GLY		3.143	57.165 137.607	1.00 20.86	0
ATOM	2202	C GLY		2.424	55.859 137.815	1.00 24.57	0
ATOM	2203	O GLY		1.365	55.615 137.223	1.00 2.00 1.00 2.00	0
MOTA	2204	N ALA		2.974 2.422	55.041 138.704 53.722 138.971	1.00 2.00	ŏ
MOTA	2206	CA ALA		3.372	52.924 139.835	1.00 2.00	Ö
MOTA	2207 2208	CB ALA		1.055	53.706 139.576	1.00 2.00	0
MOTA MOTA	2209	O ALA		0.359	52.705 139.460	1.00 2.00	0
ATOM	2210	N GLU	230	0.666	54.802 140.220	1.00 2.00 1.00 2.00	0
MOTA	2212	CA GLU		-0.648	54.881 140.875 56.193 141.647	1.00 66.07	Ö
MOTA	2213	CB GLU		-0.787	56.014 143.111	1.00 71.18	ŏ
ATOM	2214	CG GLU		-1.123 -2.376	55.175 143.337	1.00 2.00	0
MOTA	2215 2216	OE1 GLU		-2.275	53.918 143.199	1.00 2.00	0
MOTA MOTA	2217	OE2 GLU		-3.448	55.773 143.663	1.00 2.00	0
MOTA	2218	C GLU		-1.769	54.777 139.862	1.00 2.00	0
MOTA	2219	O GLU		-2.649	53.922 139.966	1.00 60.28 1.00 12.41	Ö
MOTA	2220	N VAL		-1.684	55.652 138.861 55.749 137.772	1.00 12.41 1.00 9.72	Õ
MOTA	2222	CA VAL		-2.6 4 9 -2. 18 6	56.789 136.717	1.00 2.00	Ö
MOTA	2223	CB VAL		-3.304	57.063 135.718	1.00 2.00	0
MOTA MOTA	2224 2225	CG1 VAL		-1.728	58.077 137.396	1.00 2.00	0
MOTA	2226	C VAL		-2.B16	54.393 137.102	1.00 11.88	0
MOTA	2227	O VAL		-3.937	53.966 136.849	1.00 2.00	0
ATOM	2228	N VAI	232	-1.695	53.723 136.832	1.00 2.00 1.00 2.00	0
MOTA	2230	CA VAL		-1.678	52.397 136.203 51.817 136.151	1.00 2.00	0
ATOM	2231	CB VAL		-0.2 4 5 -0.264	50.442 135.548	1.00 2.00	0
ATOM	2232	CG1 VAL	2 32	0.204			

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ATOM	2233	CG2		232	0.664	52.724 135.376	1.00 2.00 1.00 2.00	0
MOTA	2234	C	VAL	232	-2.533 -3.449	51.413 136.992 50.787 136.454	1.00 2.00	ŏ
MOTA	2235	0	VAL ALA	232 233	-2.217	51.292 138.275	1.00 38.57	Ö
MOTA	2236 2238	N CA	ALA	233	-2.920	50.395 139.173	1.00 38.57	0
MOTA MOTA	2239	CB	ALA	233	-2.297	50.487 140.542	1.00 13.20	0
ATOM	2240	c	ALA	233	-4.426	50.681 139.245	1.00 38.57	0
ATOM	2241	ō	ALA	233	-5.255	49.763 139.166	1.00 9.05	0
MOTA	2242	N	LYS	234	-4.780	51.954 139.397	1.00 2.00	0
ATOM	2244	CA	LYS	234	-6.177	52.355 139.477	1.00 2.00 1.00 22.96	0
MOTA	2245	CB	LYS	234	-6.270	53.843 139.803 54.160 141.205	1.00 22.30	Ö
MOTA	2246	cc	LYS	234	-5.776 -6.667	53.499 142.257	1.00 42.09	ŏ
MOTA	2247	CD	LYS LYS	234 234	-5.916	53.151 143.543	1.00 40.07	Ō
MOTA	2248 2249	CE NZ	LYS	234	-5.104	51.892 143.444	1.00 41.59	0
atom Atom	2253	C	LYS	234	-6.920	52.036 138.183	1.00 2.00	0
ATOM	2254	ŏ	LYS	234	-7.936	51.319 138.206	1.00 20.45	0
ATOM	2255	N	PHE	235	-6.389	52.532 137.061	1.00 14.37	0
MOTA	2257	CA	PHE	235	-6.973	52.319 135.733	1.00 14.37 1.00 2.00	0
ATOM	2258	CB	PHE	235	-6.055	52.880 134.640 52.452 133.249	1.00 2.00	ŏ
ATOM	2259	CG	PHE	235	-6. 4 38 -7.352	53.199 132.502	1.00 2.00	ŏ
MOTA	2260		PHE	235 235	-5.924	51.270 132.706	1.00 2.00	Ŏ
ATON	2261		PHE	235	-7.760	52.777 131.236	1.00 2.00	0
ATOM	2262 2263	CE2	PHE	235	-6.316	50.834 131.452	1.00 2.00	0
MOTA MOTA	2264	CZ	PHE	235	-7.242	51.588 130.710	1.00 2.00	0
ATOM	2265	C	PHE	235	-7.229	50.849 135.444	1.00 14.37	0
ATOM	2266	ō	PHE	235	-8.312	50.471 134.977	1.00 2.00 1.00 3.08	0
ATOM	2267	N	LEU	236	-6.217	50.029 135.690	1.00 3.08 1.00 5.72	Ö
ATOM	2269	CA	LEU	236	-6.339	48.606 135.447 47.900 135.731	1.00 2.00	ŏ
MOTA	2270	CB	LEU	236	-5.018 -3.915	48.136 134.710	1.00 2.00	ō
MOTA	2271	CG	LEU	236 236	-2.699	47.332 135.096	1.00 2.00	0
MOTA	227 2	CDI	LEU				1.00 2.00	0
	222	സാ		236	-4.405	47.737 133.341		
MOTA	2273	_	LEU	236 236	-4.405 -7.452	47.737 133.321 47.978 136.278	1.00 10.30	0
MOTA	2274	C	LEU LEU	236	-4.405 -7.452 -8.389	47.978 136.278 47.398 135.712	1.00 10.30 1.00 2.00	0
MOTA MOTA	2274 2275	C	LEU		-7.452 -8.389 -7.368	47.398 135.712 48.118 137.606	1.00 10.30 1.00 2.00 1.00 8.44	0
ATOM ATOM ATOM	2274	C	Leu Leu	236 236 237 237	-7.452 -8.389 -7.368 -8.368	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44	0 0 0
MOTA MOTA	2274 2275 2276	С 0 N	LEU LEU HIS HIS HIS	236 236 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56	0
MOTA MOTA MOTA MOTA MOTA MOTA	2274 2275 2276 2278 2279 2280	C O CA CB CG	LEU LEU HIS HIS HIS	236 236 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44	0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2274 2275 2276 2278 2279 2280 2281	C N CA CB CG CD2	LEU LEU HIS HIS HIS HIS	236 236 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82	0 0 0 0 0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2274 2275 2276 2278 2279 2280 2281 2282	C N CA CB CCD CD2 ND1	LEU LEU HIS HIS HIS HIS HIS HIS	236 236 237 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85	0 0 0 0 0 0 0 0
ATON ATOM MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2274 2275 2276 2278 2279 2280 2281 2282 2284	C O N CA CB CCD CD2 ND1 CE1	LEU LEU HIS HIS HIS HIS HIS HIS HIS	236 236 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26	0 0 0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285	CON CA CB CG CD2 ND1 CE1 NE2	LEU LEU HIS HIS HIS HIS HIS HIS	236 236 237 237 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44	0 0 0 0 0 0 0 0 0
ATON ATOM MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2274 2275 2276 2278 2279 2280 2281 2282 2284	C O N CA CB CCD CD2 ND1 CE1	LEU LEU HIS HIS HIS HIS HIS HIS HIS HIS HIS	236 236 237 237 237 237 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287	C O N CA CB CG CD2 ND1 CE1 NE2 C	LEU LEU HIS	236 236 237 237 237 237 237 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.355 47.031 142.350 48.033 138.138 47.232 138.027 49.346 137.947	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00	0 0 0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2285 2287 2288 2289 2291	CONCACA CONCAC	LEU LEU HIS HIS HIS HIS HIS HIS HIS HIS LYS	236 236 237 237 237 237 237 237 237 237 237 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.355 47.031 142.350 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 2.00 1.00 28.00	00000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287 2288 2291 2292	CONCACE CONCAC	LEU LEU HIS HIS HIS HIS HIS HIS HIS LYS LYS	236 236 237 237 237 237 237 237 237 237 237 238 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 2.00 1.00 29.80	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2288 2289 2291 2292 2293	CONCACBCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	LEU LEU HIS HIS HIS HIS HIS HIS LYS LYS LYS	236 236 237 237 237 237 237 237 237 237 237 238 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.927 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 2.00 1.00 28.00 1.00 28.00 1.00 28.00 1.00 36.02	00000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287 2288 2289 2291 2293 2294	CONCACBCCD2 ND1 CE1 NE2 CONCACBCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	LEU LEU HIS HIS HIS HIS HIS HIS LYS LYS LYS LYS	236 237 237 237 237 237 237 237 237 237 237	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 28.00 1.00 28.00 1.00 28.00 1.00 36.02 1.00 32.99	000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287 2288 2291 2293 2293 2294 2295	C O N CA CB CCD C O N CA CCB CCD CCD CCD CCD CCD CCD CCD CCD CCD	LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS	236 236 237 237 237 237 237 237 237 237 237 238 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.833	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 28.00 1.00 28.00 1.00 36.02 1.00 32.99 1.00 30.09	0000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287 2288 2289 2291 2293 2294	CONCACBCCD2 ND1 CE1 NE2 CONCACBCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	LEU LEU HIS HIS HIS HIS HIS HIS LYS LYS LYS LYS	236 236 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 29.80 1.00 29.80 1.00 29.80 1.00 32.99 1.00 32.99 1.00 30.09 1.00 2.00	00000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2288 2291 2292 2293 2293 2296	CON CA CBCCD2 ND1 CE1 NE2 CON CA CBCCD2 CD2 CD2 CD2 CD2 CD2 CD2 CD2 CD2 CD2	LEU LEU LEU HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS	236 236 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.827 549.215 136.403 49.205 136.234	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 2.00 1.00 29.80 1.00 36.02 1.00 32.99 1.00 30.09 1.00 2.00 1.00 2.00 1.00 2.00 1.00 30.09 1.00 2.00	0000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2289 2291 2293 2294 2295 2296 2301 2302	CON CA CB CCD CC CC CC NC CON	LEU LEU LEU HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS	236 236 237 237 237 237 237 237 237 238 238 238 238 238 238 238 238 238 238	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.843 49.215 136.403 49.205 136.234 48.607 135.584	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 38.56 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 29.80 1.00 29.80 1.00 36.02 1.00 36.02 1.00 32.99 1.00 29.21	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287 2289 2291 2292 2293 2294 2295 2296 2300 2301 2302 2304	C O N CA CB CGC CD C C C C C C C C C C C C C C C C	LEU LEU LEU HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS LYS LYS	236 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 49.26 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 2.00 1.00 29.80 1.00 36.02 1.00 36.02 1.00 30.09 1.00 29.21 1.00 29.21 1.00 2.00 1.00 29.21 1.00 2.00 1.00 29.21 1.00 2.00 1.00 29.21	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2289 2291 2293 2294 2295 2296 23001 2301 2304 2305	C O N CA CB CCD2 CD1 CE1 NE2 C O N CA CB CCD CD C CD C CD C C C C C C C C C C	LEU LEU HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS LYS	236 237 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 238 238 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340 -10.784	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 35.43 1.00 2.00 1.00 2.00 1.00 29.80 1.00 36.02 1.00 32.99 1.00 30.09 1.00 29.21 1.00 29.21 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2289 2291 2292 2293 2294 2295 2300 2300 2301 2302 2304 2305 2306	C O N CAB CCD2 CD1 CE1 NE2 C O N CAB CCD CCD CCD CCD CCD CCD CCD CCD CCD CC	LEU LEU LEU LEU LEU LEU LYS	236 237 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.827 52.103 137.104 53.628 136.83 55.925 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091 50.689 132.785	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 46.38 1.00 50.85 1.00 2.00 1.00 2.00 1.00 29.80 1.00 32.99 1.00 30.09 1.00 29.21 1.00 2.00 1.00 29.20 1.00 29.20 1.00 200 1.00 200 1.00 200 1.00 200 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2280 2281 2282 2284 2285 2287 2299 2291 2299 2299 2299 2300 23001 23002 2304 2306 2307	C O N CAB CCD2 CD1 CE1 NE2 C O N A CBC CCD N C C C C C C C C C C C C C C C C	LEU LEU LEU HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS LYS LYS LY	236 237 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282 -10.217	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.827 52.103 137.845 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.191 50.056 133.091 50.056 133.097	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 2.00 1.00 2.00 1.00 29.80 1.00 29.80 1.00 36.02 1.00 30.09 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2293 2291 2293 2294 2295 2300 2301 2302 2304 2305 2307 2308	CON CABCCC NO. CON CABCCC NO. CON CABCCC NO. CON CABCCC NO. CON CON CABCCC NO. CO	LEU LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS HIS HIS HIS HIS HIS HIS	236 237 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282 -10.217 -10.797	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.205 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091 50.689 132.785 51.046 133.397 52.225 133.284	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 2.00 1.00 2.00 1.00 29.80 1.00 29.80 1.00 35.43 1.00 29.80 1.00 29.80 1.00 29.80 1.00 29.80 1.00 29.80 1.00 29.80 1.00 29.80 1.00 200 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2289 2291 2293 2294 2295 2300 2301 2302 2304 2305 2307 2308 2310	C O N CAB CCD2 CD1 CE1 NE2 C O N A CBC CCD N C C C C C C C C C C C C C C C C	LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS LYS LYS LY	236 236 237 237 237 237 237 237 237 238 238 238 238 238 238 238 239 239 239 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282 -10.217 -11.0797 -12.052	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091 50.689 132.785 51.046 133.397 52.225 133.284 52.036 132.914	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26 1.00 2.00 1.00 2.00 1.00 29.80 1.00 36.02 1.00 36.02 1.00 32.99 1.00 30.99 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2293 2291 2293 2294 2295 2300 2301 2302 2304 2305 2307 2308	C O N CAB CCD2 NC1 O N CAB CCD2 CCD2 O N CAB CCD CCD CCD CCD CCD CCD CCD CCD CCD CC	LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS LYS LYS LY	236 237 237 237 237 237 237 237 237 238 238 238 238 238 238 238 239 239 239 239 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282 -10.217 -12.052 -10.946	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091 50.689 132.785 51.046 133.397 52.225 133.284 46.417 134.352	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 2.00 1.00 2.00 1.00 29.80 1.00 36.02 1.00 32.99 1.00 32.99 1.00 30.09 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2292 2293 2294 2295 2296 2301 2302 2304 2305 2306 2307 2308 2311 2313 2314	CON CABCCD2 ON CABCCD2 NDC CBCCD2 ON CABCCD2	LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS HIS HIS HIS HIS HIS HIS HIS HIS HIS HI	236 237 237 237 237 237 237 237 237 238 238 238 238 238 238 239 239 239 239 239 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.873 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282 -10.777 -12.052 -10.938	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091 50.689 132.785 51.046 133.397 52.225 133.284 52.036 132.914 46.417 134.352	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 2.00 1.00 2.00 1.00 29.80 1.00 29.80 1.00 36.02 1.00 36.02 1.00 39.21 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2293 2294 2295 2300 2301 2302 2304 2305 2306 2306 2307 2308 2311 2311 2311 2311	CON CAB CCD2 ON CAB CCD NC ON	LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS HIS HIS HIS HIS ASP	236 237 237 237 237 237 237 237 237 237 238 238 238 238 238 238 239 239 239 239 239 239 239 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.833 -11.776 -12.991 -10.913 -10.784 -11.125 -12.282 -10.797 -12.052 -10.946 -10.938 -10.642	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.355 47.031 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.829 54.444 138.083 55.925 137.845 49.215 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.603 133.191 50.056 133.091 50.056 133.091 50.056 133.397 52.225 133.284 52.036 132.914 46.417 134.352 45.798 133.295 44.833 135.564	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.01 1.00 46.38 1.00 50.85 1.00 49.26 1.00 2.00 1.00 2.00 1.00 29.80 1.00 36.02 1.00 32.99 1.00 30.09 1.00 2.00	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2274 2275 2276 2278 2279 2281 2282 2284 2285 2287 2292 2293 2294 2295 2296 2301 2302 2304 2305 2306 2307 2308 2311 2313 2314	CON CABCCD2 ON CABCCD2 NDC CBCCD2 ON CABCCD2	LEU LEU HIS HIS HIS HIS HIS LYS LYS LYS LYS LYS LYS LYS LYS LYS LY	236 237 237 237 237 237 237 237 237 238 238 238 238 238 238 239 239 239 239 239 239 239 239	-7.452 -8.389 -7.368 -8.368 -8.088 -9.141 -10.323 -9.035 -10.103 -10.900 -9.747 -10.672 -9.882 -11.183 -11.071 -12.427 -12.322 -11.917 -11.873 -11.776 -12.991 -10.913 -11.340 -10.784 -11.125 -12.282 -10.777 -12.052 -10.938	47.978 136.278 47.398 135.712 48.118 137.606 47.540 138.504 47.885 139.980 47.380 140.935 47.916 141.329 46.173 141.599 45.989 142.355 47.031 142.210 48.033 138.138 47.232 138.027 49.346 137.947 49.923 137.607 51.424 137.327 52.103 137.104 53.628 136.823 55.925 136.403 49.205 136.234 48.607 135.584 47.897 134.384 48.607 135.584 47.897 134.384 48.603 133.148 50.056 133.091 50.689 132.785 51.046 133.397 52.225 133.284 52.036 132.914 46.417 134.352 45.798 133.294 545.838 133.503	1.00 10.30 1.00 2.00 1.00 8.44 1.00 8.44 1.00 46.82 1.00 46.38 1.00 50.85 1.00 49.26 1.00 8.44 1.00 2.00 1.00 2.00 1.00 29.80 1.00 29.80 1.00 36.02 1.00 36.02 1.00 39.21 1.00 2.00	

ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2319 2320 2321 2322 2323 2324 2326 2327 2328 2329 2330 2331	С	ASP ASP LEU LEU LEU LEU LEU LEU	240 240 240 240 241 241 241 241 241 241	-12.676 -12.687 -13.610 -9.393 -9.626 -8.417 -7.462 -7.086 -8.185 -7.795 -8.435 -6.225	43.919 134.87 43.571 133.66 44.578 135.39 43.985 134.38 42.933 133.79 44.823 134.04 44.551 132.95 45.858 132.22 46.590 131.42 48.047 131.22 45.899 130.08 43.925 133.60 43.832 134.81	6 1.00 50.73 6 1.00 47.58 5 1.00 37.76 7 1.00 40.81 0 1.00 2.00 5 1.00 2.00 9 1.00 2.00 9 1.00 2.00 4 1.00 2.00 7 1.00 2.00 1 1.00 2.00	000000000000
MOTA	2332 2333 2335	O N CA	ASP ASP	241 242 242	-5.235 -4.046	43.534 132.81 42.895 133.38	7 1.00 2.00	0
MOTA MOTA	2336	CB	ASP	242	-3.974	41.455 132.91 40.511 133.82	5 1.00 25.69	0
MOTA MOTA	2337 2338	CG OD1	ASP ASP	242 242	-4.683 - 5.751	40.853 134.36		Ö
MOTA	2339	OD2	ASP	242	-4.149	39.404 133.97		0
MOTA MOTA	2340 2341	C O	ASP ASP	242 242	-2.711 -1.702	43.506 133.03 43.214 133.67		0
ATOM	2342	N	LEU	243	-2.689	44.317 131.99	4 1.00 23.97	0
MOTA MOTA	2344 2345	CA CB	LEU	243 243	-1.440 -0.644	44.899 131.53 43.799 130.82		0
ATOM	2346	œ	LEU	243	0.698	43.831 130.09	7 1.00 2.00	0
MOTA	2347	-	LEU	243 243	0.410 1.526	43.690 128.66 45.058 130.39		0
MOTA MOTA	2348 2349	CD2	LEU	243	-1.804	46.013 130.59	0 1.00 19.91	0
MOTA	2350	0	LEU	243	-2.960	46.149 130.18 46.849 130.28		0
MOTA MOTA	2351 2353	N CA	ILE	244 244	-0.834 -1.070	47.919 129.35	4 1.00 22.40	0
ATOM	2354	СВ	ILE	244	-1.164	49.269 130.06		0
MOTA	2355	CG2	ILE	244 244	-1. 23 5 -2. 39 5	50.405 129.02 49.255 130.99		0
MOTA MOTA	2356 2357	CG1 CD1	ILE	244	-2.607	50.530 131.80	3 1.00 2.00	0
MOTA	2358	С	ILE	244	0.069	47.902 128.36		0
MOTA MOTA	2359 2 36 0	O N	CYS	244 245	1.234 -0.286	47.715 128.75 48.063 127.09		0
ATOM	2362	CA	CYS	245	0.675	48.074 126.01	9 1.00 2.00	0
MOTA	2363	CB	CYS	245	0.403. 1.849	46.930 125.07 45.913 124.98		0
ATOM ATOM	2364 2365	SG C	CYS	245 245	0.668	49.389 125.27		0
ATOM	2366	ō	CYS	245	-0.362	49.842 124.78	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	0
MOTA	2367	N	ARG	246 246	1.829 1.987	50.014 125.23 51.277 124.56		0
ATOM ATOM	2369 2370	CA CB	ARG	246	1.763	52.438 125.54	1 1.00 2.00	0
MOTA	2371	CG	ARG	246	2.658	52.453 126.78		0
MOTA MOTA	2372 2373	CD NE	ARG ARG	246 246	3.969 3.714	53.245 126.62 54.656 126.38		Ö
MOTA	2375	CZ	ARG	246	4.610	55.622 126.51		0
ATOM	2376		ARG ARG	246 246	5.842 4.263	55.334 126.89 56.881 126.28		0
MOTA MOTA	2379 2382	C	ARG	246	3.374	51.342 123.93	8 1.00 2.00	0
MOTA	2383	0	ARG	246	4.216	50.471 124.16 52.345 123.10		0
MOTA MOTA	2384 2386	N CA	ALA ALA	247 · 247	3.605 4.906	52.489 122.48		Ŏ
ATOM	2387	СВ	AL A	247	4.791	52.272 120.97		0
ATOM ATOM	2388	C	ALA ALA	247 247	5.351 5.806	53.902 122.819 54.172 123.92		0
ATOM	2389 2390	O N	HIS	248	5.187	54.801 121.87	0 1.00 2.00	0
MOTA	2392	CA	HIS	248	5.512	56.193 122.04		0
MOTA MOTA	2393 2394	С 0	HIS HIS	248 248	6.959 7.507	56.579 122.36 57.453 121.67	2 1.00 2.00	0
ATOM	2395	CB	HIS	248	4.539	56.793 123.05	4 1.00 2.00	0
MOTA	2396	CG	HIS	248	4.255 5.262	58.246 122.83° 59.147 122.59		0
MOTA MOTA	2397 2398		HIS	248 248	4.677	60.327 122.53	5 1.00 2.0 0	0
MOTA	2399		HIS	248	3.074	58.906 122.91	3 1.00 2.00	0

					2.250	60.234 122.	721	1.00	2.00	0
MOTA	2400	NE2		248	3.359 7.588	55.942 123.3	348		7.57	ŏ
MOTA	2402		GLN	2 4 9 2 4 9	8.967	56.283 123.	731		7.57	0
ATOM	2404 2405	-	GLN GLN	249	9.010	56,722 125.3	192	1.00 3	0.82	0
MOTA MOTA	2406		GLN	249	7.987	57.759 125.			2.63	0
ATOM	2407		GLN	249	8.159	58.253 126.			3.33	0
ATOM	2408		GLN	249	7.871	59.406 127.2 57.389 127.1			9.63 2.74	0
MOTA	2409		GLN	249	8.640 10.052	57.389 127.0 55.222 123.9			7.57	ŏ
MOTA	2412	-	GLN	249	9.884	54.057 123.			8.32	ō
ATOM	2413	_	GLN	249 250	11.185	55.664 122.			0.32	0
MOTA	2414 2416		VAL VAL	250	12.339	54.805 122.	735		0.32	0
MOTA MOTA	2417		VAL	250	13.571	55.628 122.3	300		6.81	0
ATOM	2418	CG1		250	14.641	54.714 121.	706		6.81	0
ATOM	2419	CG2	VAL	250	13.161	56.708 121.			6.81 0.32	0
MOTA	2420	_	VXL	250	12.733 12.471	54.034 123.5 54.474 125.5			6.81	ŏ
MOTA	2421	_	VAL	250 251	13.372	52.887 123.	797		2.00	ō
MOTA	2422		VAL	251 251	13.820	52.023 124.	B71		2.00	0
MOTA	2424 2425		VAL	251	12.655	51.150 125.3	391		9.99	0
MOTA MOTA	2426	CG1		251	11.778	51.967 126.	311		9.99	0
ATOM	2427	CG2	VAL	251	11.803	50.661 124.	247		9. 9 9 2.00	0
MOTA	2428	С	VAL	251	14.960	51.167 124.5 50.470 123.5	310		9.99	ŏ
ATOM	2429	0	VAL	251	14.792 16.118	51.224 124.	972		1.55	Ō
MOTA	2430	N	GLU GLU	252 252	17.312	50.509 124.	519	1.00 1	9.34	0
ATOM ATOM	2432 2433	CA CB	GLU	252	18.452	50.605 125.	542		1.04	0
ATOM	2434	œ	GLU	252	19.819	50.220 124.	933		8.95	0
MOTA	2435	CD	GLU	252	20.926	49.966 125. 49.303 125.	962 597		0.82 7.37	Ö
MOTA	2436	OE1		252	21.938 20.782	50.422 127.			9.06	0
MOTA	2437	OE2	GLU	252 252	17.144	49.055 124.		1.00 1	9.22	0
MOTA	2438 2439	C O	GLU GLU	252	17.838	48.571 123.			8.64	Ŏ
MOTA MOTA	2440	N	ASP	253	16.240	48.356 124.			2.00 2.00	0
ATOM	2442	CA	ASP	253	16.025	46.936 124. 46.144 125.	544 950		B.16	Õ
MOTA	2443	CB	ASP	253	16.125 17.543	46.106 126.	401		4.05	0
ATOM	2444	CG	asp asp	253 253	18.060	44,995 126.	635	1.00 8	9.15	0
ATOM ATOM	2445 2446		ASP	253	18.148	47.1B1 126.			7.62	0
ATOM	2447	C	ASP	253	14.710	46.645 123.	836	1.00 1.00 6	2.00 3.68	ŏ
ATOM	2448	0	ASP	253	14.201	45.528 123. 47.660 123.		1.00	2.00	Ö
MOTA	2449	N	GLY	254	14.161 12.919	47.483 122.	457	1.00	2.00	0
MOTA	2451	CA	GLY GLY	254 254	11.715	47.547 123.		1.00	2.00	0
MOTA	2452 2453	CO	GLY	254	10.694	48.124 122.		1.00 4		0
MOTA MOTA	2454	N	TYR	255	11.823	46.922 124.		1.00	2.00	0
ATOM	2456	CA	TYR	255	10.749	46.956 125. 45.614 125.		1.00 1.00	2.00	ŏ
ATOM	2457	CB	TYR	255	10.047 10.853	45.614 125. 44.510 126.		1.00	2.00	0
ATOM	2458	CG	TYR	255 255	12.103	44.175 125.		1.00	2.00	0
MOTA	2459	CD1	TYR	255	12.824	43.121 126.	248	1.00	2.00	0
MOTA MOTA	2460 2461	CD2		255	10.337	43.770 127.	246	1.00	2.00 2.00	0
ATOM	2462	CE2		255	11.041	42.721 127	802	1.00 1.00	2.00	0
ATOM	2463	CZ	TYR	255	12.279	42.397 127 41.328 127		1.00	2.00	Õ
MOTA	2464	OH	TYR	255	12.946 11.262	47.375 126		1.00	2.00	0
MOTA	2466	C	TYR	255 255	12.453	47.283 127	.150	1.00	2.00	0
MOTA	2467	O N	TYR GLU	256	10.336	47.853 127	. 6 56	1.00	2.00	0
ATOM	2468 2470	N CA	GLU	256	10.632	48.341 128	. 9 79	1.00	2.00	0
MOTA MOTA	2471	CB	GLU	256	11.092	49.798 128	.910		20.70 27.98	0
MOTA	2472	ČĞ	GLU	256	11.264	50.457 130 51.961 130	179		31.28	ő
MOTA	2473	CD	GLU	256	11.485 10.726	52.725 130			33.21	0
MOTA	2474	OE:		256 256	12.418	52.378 129	.453	1.00	39.87	0
MOTA	2475 2476	OE:	2 GLU GLU	25 6	9.362	48.238 129	.800	1.00	2.00	0
MOTA MOTA			920			40 743 130	470	1.00	18.74	0
			GLU	256	8.299	48.743 129	. 420			U
	2477 2478	0	GLU PHE	257	9.474	47.540 130	.919	1.00	2.00	0
MOTA MOTA	2477	0 N	PHE			48.743 129 47.540 130 47.364 131	.919			

2.00 46.139 132.661 1.00 8.578 8.222 2481 CB PHE 257 MOTA 2.00 1.00 44.865 131.978 257 PHE 2482 CG MOTA 2.00 43.926 131.694 1.00 257 9.187 CD1 PHE MOTA 24B3 2.00 44.594 131.649 1.00 6.906 257 2484 CD2 PHE MOTA 2.00 1.00 42.740 131.101 257 8.849 CE1 PHE MOTA 2485 43.417 131.058 42.481 130.781 1.00 2.00 257 6.560 CE2 PHE 2486 MOTA 1.00 2.00 257 7.533 CZ PHE 2487 MOTA 48.593 132.671 1.00 2.00 8.257 PHE 257 MOTA 2488 С 2.00 49.360 132.782 1.00 0 9.215 257 2489 0 PHE MOTA 48.777 133.286 49.916 134.147 1.00 7.104 258 PHE 2490 N MOTA 1.00 25.02 258 258 258 6.884 CA PHE 2492 MOTA 2.00 1.00 51.086 133.299 0 6.417 2493 CB PHE MOTA 52.237 134.072 53.325 134.431 1.00 0 5.831 PHE CG MOTA 2494 2.00 1.00 258 258 6.625 MOTA 2495 CD1 PHE 2.00 4.468 1.00 ۵ 52.271 134.369 CD2 PHE 2496 MOTA 54.439 135.070 53.369 135.003 1.00 0 258 CE1 PHE MOTA 2497 2.00 1.00 0 258 3.900 2498 CE2 PHE MOTA 54.461 135.356 1.00 2.00 0 4.698 258 PHE CZ 2499 MOTA 5.837 4.965 5.953 49.508 135.160 48.675 134.859 1.00 25.02 1.00 2.00 258 258 PHE MOTA 2500 C 0 2501 0 PHE MOTA 50.063 136.368 1.00 13.61 259 ALA MOTA 2502 N 1.00 11.22 0 49.775 137.457 5.027 259 2504 CA ALA MOTA 50.318 137.122 48.286 137.729 2.00 1.00 0 3.626 ALA 259 CB 2505 MOTA 1.00 12.73 259 259 4.951 ALA MOTA 2506 С 47.718 137.660 2.00 0 1.00 3.876 0 ALA 2507 MOTA 1.00 2.00 0 47.657 138.016 6.091 260 LYS MOTA 2508 N 6.140 5.**59**4 1.00 9.39 0 46.214 138.304 LYS 260 MOTA 2510 CA LYS 1.00 20.75 0 45.925 139.704 2511 CB 260 MOTA 46.151 140.844 47.622 141.177 47.765 142.449 6.589 6.797 1.00 28.17 260 LYS MOTA 2512 CG 1.00 36.91 0 260 2513 CD LYS MOTA 1.00 36.02 0 LYS 7.615 260 2514 CE MOTA 6.919 1.00 29.33 47.158 143.633 45.375 137.279 0 260 2515 2519 NZ LYS MOTA 2.00 1.00 0 5.370 LYS 260 С ATOM 44.633 137.641 1.00 15.95 4.443 5.764 260 2520 2521 2.00 MOTA 0 LYS 1.00 0 45.531 136.004 ATOM ARG 261 N 0 44.843 134.851 1.00 2.00 ARG 5.173 261 CA 2523 MOTA 43.335 134.964 1.00 21.35 5.410 6.881 261 CB ARG 2524 MOTA 1.00 21.35 0 42.969 134.928 2525 ARG 261 MOTA CG 41.473 134.970 41.097 134.460 1.00 9.03 0 7.088 261 2526 ARG CD MOTA 8.408 8.801 1.00 14.18 261 2527 NE ARG MOTA 0 1.00 14.38 39.842 134.222 ARG 261 2529 CZ MOTA 38.820 134.462 39.603 133.729 1.00 16.12 0 7.976 10.018 261 MOTA 2530 NH1 ARG 1.00 18.11 2533 NH2 ARG 261 MOTA 45.136 134.592 n 1.00 2.00 3.692 261 2536 ARG MOTA С 44.343 133.933 46.283 135.092 46.727 134.927 1.00 21.35 1.00 7.73 0 261 3.009 2537 0 ARG MOTA 3.215 2538 GLN 262 N MOTA 1.00 7.73 0 1.821 GLN 262 MOTA 2540 CA 2.00 47.620 136.100 1.00 1.379 MOTA 2541 GLN 262 CR 46.888 137.434 47.840 138.601 2.00 0 1.00 1.106 2542 GLN 262 CG MOTA 1.00 2.00 262 0.859 GLN MOTA 2543 CD 1.00 2.00 -0.277 48.137 138.930 ATOM 2544 OE1 GLN 262 48.304 139.237 47.495 133.628 2.00 1.00 1.922 2545 262 NE2 GLN MOTA 1.00 7.73 1.635 262 MOTA 2548 GLN 2.00 0 1.00 0.526 47.650 133.154 GLN 262 2549 0 MOTA 47.996 133.078 48.726 131.819 1.00 2.00 2.730 263 LEU MOTA 2550 N 1.00 0 2.00 2.723 2.754 263 2552 CA LEU MOTA 2.00 50.244 132.069 1.00 CB LEU 263 2553 MOTA 51.201 130.890 1.00 2.00 1.00 2.00 0 3.070 263 MOTA 2554 CG LEU 52.532 131.133 2.404 CD1 LEU 263 2555 MOTA 51.408 130.679 1.00 2.00 4.575 CD2 LEU 263 MOTA 2556 1.00 2.00 3.991 48.305 131.089 MOTA 2557 LEU 263 С 48.082 131.736 Ω 1.00 2.00 263 5.018 2558 LEU 0 MOTA 48.207 129.759 1.00 42.92 3.932 264 MOTA 2559 N VAL 1.00 41.59 5.105 47.850 128.949 ATOM VAL 264 CA 2561 46.373 128.433 2.00 1.00 264 5.014 VAL MOTA 2562 CB 46.242 127.372 1.00 2.00 3.952 CG1 VAL 264 MOTA 2563 6.**34**9 5.**19**3 45.908 127.929 1.00 2.00 CG2 VAL 264 2564 MOTA 1.00 42.79 48.840 127.772 264 MOTA 2565 VAL

ATOM	2566	0	VAL	264	4.164	49.261	127.242	1.00	2.00	0
ATOM	2567	N	THR	265	6.409		127.400	1.00	2.00	Ŏ
MOTA	2569	CA	THR	265	6.639	50.177	126.292	1.00	2.00	0
MOTA	2570	CB	THR	265	7.420		126.789	1.00	2.00	0
MOTA	2571	OG1	THR	265	6.534		127.531	1.00	2.00	0
MOTA	2573	CG2	THR	265	8.021		125.648	1.00	2.00	0
MOTA	2574	Č	THR	265	7.405		125.132	1.00	2.00	0
MOTA	2575	0	THR	265	8.612 6.712		125.246 124.019	1.00	2.00 7.19	0
MOTA MOTA	2576	N	LEU	266 266	7.330		122.857	1.00	7.19	0
ATOM	2578 2579	CA CB	LEU	266	6.338		122.176	1.00	2.00	Ö
ATOM	2580	CG	LEU	266	5.815		122.948	1.00	2.00	ŏ
ATOM	2581		LEU	266	4.859		122.057	1.00	2.00	ŏ
MOTA	2582		LEU	266	6.955	45.599	123.368	1.00	2.00	ō
MOTA	2583	C	LEU	266	7.898		121.819	1.00	7.19	Ó
MOTA	2584	0	LEU	26 6	7.329		121.537	1.00	2.00	0
MOTA	2585	N	PHE	267	9.033		121.247	1.00	2.00	0
MOTA	2587	CA	PHE	267	9.665		120.222	1.00	2.00	0
ATOM	2588	CB	PHE	267	10.763		120.821	1.00	2.00	0
ATOM	2589	CG	PHE	267	10.937		120.091	1.00	2.00	0
MOTA	2590		PHE	267	9.985		120.206	1.00	2.00	0
ATOM	2591	CD2		267	12.021		119.255 119.493	1.00	2.00	0
MOTA	2592	CE1		267	10.108 12.146		118.540	1.00	2.00 2.00	0
MOTA	2593	CE2	PHE	267 267	11.187		118.661	1.00	2.00	ŏ
MOTA MOTA	2594 2595	CZ C	PHE PHE	267	10.246		119.227	1.00	2.00	ŏ
ATOM	2596	Ö	PHE	267	11.418		119.321	1.00	2.00	ŏ
ATOM	2597	N	SER	268	9.387		118.302		12.94	ō
ATOM	2599	CA	SER	268	9.707		117.267	1.00	12.94	0
ATOM	2600	CB	SER	268	8.420	47.043	116.723	1.00	11.11	0
MOTA	2601	OG	SER	268	7.593		117.771		11.11	0
ATOM	2603	С	SER	268	10.450		116.10 3		12.94	0
MOTA	2604	0	SER	268	10.150		115.711		11.11	0
MOTA	2605	N	ALA	269	11.391			1.00		0
MOTA	2607	CA	ALA	269	12.222		114.378		71.72	0
MOTA	2608	CB	ALA	269	11.383		113.295		82.00 74.82	0
MOTA	2609	C	ALA	269	13.585 14.609		114.543 114.186		91.21	Ö
MOTA MOTA	2610 2611	O N	ALA PRO	269 270	13.619		115.076		28.82	ŏ
ATOM	2612	CD	PRO	270	12.436		115.513	1.00	2.00	ŏ
ATOM	2613	CA	PRO	270	14.788		115.316	1.00	31.86	0
MOTA	2614	CB	PRO	270	14.340	51.374	116.502	1.00	2.00	0
MOTA	2615	CG	PRO	270	13.034		116.023	1.00	2.00	0
MOTA	2616	С	PRO	270	16.254		115.428		29.52	0
MOTA	2617	0	PRO	270	16.716		115.272	1.00	2.00	0
ATOM	2618	N	asn	271	16.928		115.593	1.00	2.00	0
MOTA	2620	CA	asn	271	18.346		115.754	1.00	2.00	0
MOTA	2621	CB	ASN	271	19.168		114.663		35.96 61.96	0
MOTA	2622	CG	ASN	271	18.483		113.300 113.034		36.11	ő
MOTA	2623	OD1		271 271	17.605 18.872		112.432		36.31	ŏ
ATOM ATOM	2624 2627	ND2 C	ASN	271	18.199	53.143	115.475	1.00	2.00	0
ATOM	2628	0	ASN	271	19.072	53.B07	114.915		61.69	0
ATOM	2629	N	TYR	272	17.039		115.930	1.00	2.00	0
ATOM	2631	CA	TYR	272	16.489	54.978	115.793	1.00	2.00	0
MOTA	2632	CB	TYR	272	15.772	55.358	117.090		17.3 3	0
ATOM	2633	CG	TYR	272	14.528	56.208	116.897		10.41	0
MOTA	2634	CD1		272	13.458		116.105		11.86	0
ATOM	2635	CE1	TYR	272	12.313		115.917		14.32	0
MOTA	2636	CD2	TYR	272	14.421		117.493		10.66 15.43	0
ATOM	2637	CE2	TYR	272	13.287		117.312 116.529	1.00	9.40	0
MOTA	263B	CZ	TYR	272 2 7 2	12.243 11.143	58 636	116.375		17.38	ŏ
MOTA	2639	C OH	TYR TYR	272	17.332		115.299	1.00	2.00	Ö
ATOM ATOM	2641 2642	0	TYR	272	18.356		115.888		29.84	Ö
ATOM	2643	N	CYS	273	16.860	56.731	114.201	1.00	7.24	0
ATOM	2645	CA	CYS	273	17.490		113.553	1.00	6.82	0
ATOM	2646	CB	CYS	273	17.203		114.331	1.00	11.61	0

MOTA MOTA MOTA MOTA	2647 2648 2649 2650	SG C O N	CYS CYS CYS GLY	273 273 273 274	15.496 19.010 19.738 19.498	57.724 58.717 56.502	114.843 113.411 113.512 113.185	1.00 10.42 1.00 8.43 1.00 9.04 1.00 13.82	0 0 0
MOTA	2652	CA	GLY	274	20.934		113.044	1.00 13.82 1.00 13.82	0
ATOM	2653	С	GLY	274	21.710		114.134 113.906	1.00 13.82	0
MOTA	2654	0	GLY	2 74 275	22.817 21.120	57.085	115.326	1.00 92.80	Ö
MOTA	2655 2657	N CA	GLU GLU	275	21.719	57.764	116.467	1.00 91.32	0
MOTA MOTA	2658	CB	GLU	275	21.479	59.276	116.340	1.00 25.01	0
ATOM	2659	ĊĠ	GLU	275	19.994		116.224	1.00 31.82 1.00 30.92	0
MOTA	2660	CD	GLU	275	19.730 18.951		116.060 116.868	1.00 30.92 1.00 38.31	0
MOTA	2661	OE1 OE2	GLU GLU	275 275	20.276		115.125	1.00 34.18	ŏ
MOTA MOTA	2662 2663	C	GLU	275	21.137	57.260	117.794	1.00 90.32	0
MOTA	2664	ō	GLU	275	20.941		118.716	1.00 23.69	0
MOTA	2665	N	PHE	276	20.868	55.958	117.901 119.134	1.00 18.78 1.00 17.07	0
ATOM	2667	CA	PHE	276 276	20.303 18.774	55.551	119.125	1.00 37.20	ŏ
MOTA MOTA	2668 2669	CB CG	PHE	276	18.280	56.B76	119.652	1.00 35.89	0
ATOM	2670	CD1		276	17.522	57.726	118.849	1.00 35.53	0
MOTA	2671	CD2	PHE	276	18.572	57.273	120.949	1.00 39.17 1.00 31.45	0
MOTA	2672		PHE	276	17.067 18.123	58.485	119.329 121.434	1.00 31.43	0
MOTA	2673	CE2 CZ	PHE PHE	276 276	17.367	59.324	120.621	1.00 39.75	ō
MOTA MOTA	2674 2675	C	PHE	276	20.681	53.992	119.597	1.00 17.79	0
MOTA	2676	ō	PHE	276	20.463	53.669	120.772	1.00 38.38	0
MOTA	2677	N	ASP	277	21.213	53.140	118.712 119.093	1.00 27.69 1.00 27.89	0
MOTA	2679	CA	ASP	277 277	21.638 22.884	51.772 51.848	120.018	1.00 0.27	ŏ
MOTA	2680 2681	CB CG	ASP ASP	277	23.231	50.508	120.716	1.00 39.20	0
MOTA MOTA	2682		ASP	277	23.163	50.470	121.964	1.00 39.20	0
MOTA	2683	OD2	ASP	277	23.576		120.036	1.00 39.20 1.00 26.41	0
MOTA	2684	C	ASP	277	20.560		119.742 119.918	1.00 26.41 1.00 0.76	ŏ
MOTA	2685	0	asp asn	277 278	20.761 19.428	51.491		1.00 2.00	ŏ
MOTA MOTA	2686 2688	N CA	ASN	278	18.344	50.758	120.722	1.00 2.00	0
MOTA	2689	CB	ASN	278	17.142		120.920	1.00 2.00 1.00 2.00	0
MOTA	2690	CG	ASN	278	16.394 16.867	51.930 52.643	119.639 118.765	1.00 2.00	ŏ
MOTA	2691 2692	ND2	asn asn	278 278	15.225	51.324		1.00 2.00	ō
MOTA MOTA	2695	C	ASN	278	17.938		119.829	1.00 2.00	0
ATOM	2696	ō	ASN	278	17.970	49.710	118.600	1.00 2.00	0
MOTA	2697	N	ALA	279				1 00 14 71	Λ
MOTA	2699				17.603		120.454	1.00 14.31	0
MOTA		CA	ALA	279	17.133	47.320	119.734	1.00 14.31 1.00 2.00	0
MOTE	2700	CB	ALA	279 279		47.320	120.454 119.734 120.594 119.498	1.00 14.31 1.00 2.00 1.00 14.31	0
MOTA MOTA				279	17.133 17.281 15.655 15.155	47.320 46.081 47.631 48.645	119.734 120.594 119.498 119.996	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00	0 0 0
MOTA MOTA	2700 2701 2702 2703	CB C O N	ALA ALA ALA GLY	279 279 279 279 280	17.133 17.281 15.655 15.155 14.959	47.320 46.081 47.631 48.645 46.788	119.734 120.594 119.498 119.996 118.743	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00	0 0 0
MOTA MOTA	2700 2701 2702 2703 2705	CB C O N CA	ALA ALA ALA GLY GLY	279 279 279 279 280 280	17.133 17.281 15.655 15.155 14.959 13.549	47.320 46.081 47.631 48.645 46.788 47.014	119.734 120.594 119.498 119.996 118.743 118.484	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00	0 0 0
ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706	CB C O N CA C	ALA ALA ALA GLY GLY GLY	279 279 279 279 280 280 280	17.133 17.281 15.655 15.155 14.959 13.549 12.864	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698	119.734 120.594 119.498 119.996 118.743 118.484 118.992 118.799	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 9.35	0 0 0 0 0 0
MOTA MOTA ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707	CB C O N CA C	ALA ALA ALA GLY GLY	279 279 279 279 280 280	17.133 17.281 15.655 15.155 14.959 13.549	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906	119.734 120.594 119.498 119.996 118.743 118.484 118.992 118.799 119.643	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 9.35 1.00 2.00	00000000
ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2710	CB C O N CA C	ALA ALA GLY GLY GLY GLY ALA ALA	279 279 279 279 280 280 280 281 281	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.725 11.082	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 44.702	119.734 120.594 119.498 119.996 118.743 118.484 118.992 118.799 119.643 120.167	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 9.35 1.00 2.00 1.00 2.00	000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711	CB C O N CA C O N CA CB	ALA ALA GLY GLY GLY GLY ALA ALA	279 279 279 279 280 280 280 281 281 281	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.725 11.082 10.799	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.702 44.850	119.734 120.594 119.498 119.996 118.743 118.484 118.992 118.799 119.643 120.167 121.658	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 9.35 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	00000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711 2712	CB CONCA CONCA CBC	ALA ALA GLY GLY GLY ALA ALA ALA	279 279 279 279 280 280 280 281 281 281	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.725 11.082 10.799 9.815	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.702 44.850 44.345	119.734 120.594 119.498 119.996 118.743 118.484 118.992 119.643 120.167 121.658 119.437	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 9.35 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711 2712 2713	CB CONCA CONCA CB COO	ALA ALA GLY GLY GLY ALA ALA ALA ALA	279 279 279 280 280 280 281 281 281 281 281	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.725 11.082 10.799	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.702 44.850 44.345 44.345 44.345	119.734 120.594 119.498 119.996 118.743 118.484 118.992 119.643 120.167 121.658 119.437 119.918	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 14.19	00000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711 2712	CB CONCA CONCA CBC	ALA ALA GLY GLY GLY ALA ALA ALA	279 279 279 279 280 280 280 281 281 281 281 282 282	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.725 11.082 10.799 9.815 9.451 9.140 7.907	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.702 44.850 44.345 44.992 43.312 42.833	119.734 120.594 119.498 119.996 118.743 118.484 118.992 119.643 120.167 121.658 119.437 118.473 119.918	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 14.19 1.00 14.19	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711 2712 2713 2714 2716 2717	CB CONCACONCACBCONCACCBCCCCCCCCCCCCCCCCCCCC	ALA ALA GLY GLY GLY ALA ALA ALA ALA MET MET MET	279 279 279 279 280 280 281 281 281 281 282 282	17.133 17.281 15.655 15.155 14.959 13.381 11.725 11.082 10.799 9.815 9.451 9.140 7.907 8.232	47.320 46.081 47.631 48.645 46.788 47.018 45.906 44.702 44.850 44.345 44.992 43.312 42.815	119.734 120.594 119.498 119.996 118.743 118.799 119.643 120.167 121.658 119.437 118.473 119.918 119.918	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.19 1.00 14.19 1.00 25.98	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711 2712 2713 2714 2716 2717 2718	CB CONCA CONCA CCONCA CCCONCA CCCCC	ALA ALA GLY GLY GLY ALA ALA ALA MET MET MET	279 279 279 280 280 280 281 281 281 281 282 282 282	17.133 17.281 15.655 15.155 14.959 13.381 11.725 11.082 10.799 9.815 9.451 9.140 7.907 8.232 7.056	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.996 44.702 44.850 44.345 44.992 43.312 42.833 42.115 40.449	119.734 120.594 119.498 119.996 118.743 118.484 118.992 119.643 120.167 121.658 119.437 118.473 119.918 119.315 117.060 115.980	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 14.19 1.00 14.19	000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2710 2711 2712 2713 2714 2716 2717 2718 2719	CB CONCA CONCA CC CONCA CC CC CC CC CC CC CC CC CC CC CC CC C	ALA ALA GLY GLY GLY ALA ALA ALA ALA MET MET MET MET	279 279 279 279 280 280 281 281 281 281 282 282	17.133 17.281 15.655 15.155 14.959 13.381 11.725 11.082 10.799 9.815 9.451 9.140 7.907 8.232	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 44.702 44.850 44.345 44.992 43.312 42.833 42.115 40.449 40.644	119.734 120.594 119.498 119.996 118.743 118.484 118.992 118.658 119.437 118.473 119.315 117.960 115.980 115.525	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.19 1.00 14.19 1.00 24.91 1.00 25.45 1.00 21.20	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2710 2711 2712 2713 2714 2716 2717 2718	CB CONCA CONCA CCONCA CCCONCA CCCCC	ALA ALA GLY GLY GLY ALA ALA ALA MET MET MET	279 279 279 280 280 281 281 281 281 282 282 282 282	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.725 11.082 10.799 9.815 9.451 9.140 7.907 8.232 7.056 7.304 9.044 7.397	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.702 44.850 44.345 44.345 42.115 41.885 40.449 40.644 41.851	119.734 120.594 119.996 119.996 118.743 118.484 118.992 119.643 120.167 121.658 119.437 119.918 119.315 117.960 115.980 115.525 120.363	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.19 1.00 14.19 1.00 25.45 1.00 21.20 1.00 14.19	000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2711 2712 2713 2714 2716 2717 2718 2719 2720 2721 2722	C C O N C C O O N C C C C O O N C C C C	ALA ALA GLY GLY GLY ALA ALA ALA ALA MET MET MET MET MET MET MET MET	279 279 279 280 280 281 281 281 281 282 282 282 282 282 282	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.082 10.799 9.815 9.451 9.140 7.907 8.232 7.056 7.304 9.044 7.397 8.142	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.345 44.850 44.345 41.885 40.449 40.644 41.851 40.972	119.734 120.594 119.996 119.996 118.743 118.484 118.992 118.799 119.643 120.167 121.658 119.437 119.918 119.315 117.995 117.060 115.980 115.525 120.363 120.790	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.19 1.00 14.19 1.00 25.98 1.00 24.91 1.00 24.98	000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2711 2712 2713 2714 2716 2717 2718 2719 2720 2721 2722 2723	CB CONCOONCCOONCCCSCOON	ALA ALA GLY GLY GLY ALA ALA ALA ALA MET MET MET MET MET MET MET MET MET	279 279 279 279 280 280 281 281 281 282 282 282 282 282 282 282	17.133 17.281 15.655 15.155 14.959 13.381 11.725 11.082 10.799 9.815 9.845 19.140 7.907 8.232 7.056 7.304 9.044 7.397 8.142 6.148	47.320 46.081 47.631 48.645 46.788 47.014 45.783 44.698 45.906 44.702 44.850 44.992 42.833 42.115 41.885 40.449 40.644 41.8572 42.004	119.734 120.594 119.498 119.996 118.743 118.484 118.992 119.643 120.167 121.658 119.437 118.473 119.315 117.995 117.060 115.525 120.363 120.798	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.19 1.00 14.19 1.00 25.45 1.00 21.20 1.00 14.19	000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2700 2701 2702 2703 2705 2706 2707 2708 2711 2712 2713 2714 2716 2717 2718 2719 2720 2721 2722	C C O N C C O O N C C C C O O N C C C C	ALA ALA GLY GLY GLY ALA ALA ALA ALA MET MET MET MET MET MET MET MET	279 279 279 280 280 281 281 281 281 282 282 282 282 282 282	17.133 17.281 15.655 15.155 14.959 13.549 12.864 13.381 11.082 10.799 9.815 9.451 9.140 7.907 8.232 7.056 7.304 9.044 7.397 8.142	47.320 46.081 47.631 48.645 46.788 47.0783 44.698 45.906 44.702 44.850 44.345 44.992 43.312 42.833 41.885 40.449 40.644 41.851 40.972 42.004 41.143	119.734 120.594 119.996 119.996 118.743 118.484 118.992 118.799 119.643 120.167 121.658 119.437 119.918 119.315 117.995 117.060 115.980 115.525 120.363 120.790	1.00 14.31 1.00 2.00 1.00 14.31 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 14.19 1.00 14.19 1.00 25.98 1.00 24.91 1.00 25.45 1.00 21.20 1.00 24.98 1.00 24.98 1.00 24.98	000000000000000000000000000000000000000

ATOM	2727	CG	MET	283	4.481	41.205	124.142	1.00 19.76	0
MOTA	2728	SD	MET	283	3.228	42.077	125.116	1.00 20.92	0
MOTA	2729	CE	MET	283	4.204		125.966	1.00 17.38 1.00 2.00	0
MOTA	2730	C	MET	283	4.592		121.339 121.036	1.00 2.00	0
MOTA	2731	0	MET SER	283 284	3. 45 6 5. 012		121.291	1.00 2.00	ŏ
MOTA MOTA	2732 2734	N CA	SER	284	4.148	37.777	120.869	1.00 2.00	ŏ
ATOM	2735	CB	SER	284	4.991	36.664	120.220	1.00 20.90	Ö
ATOM	2736	OG	SER	284	6.376	36.771	120.528	1.00 23.67	0
ATOM	2738	C	SER	284	3.275		122.018	1.00 2.00	0
MOTA	2739	0	SER	284	3.777		122.978	1.00 26.24	0
MOTA	2740	N	VAL	285	1.967		121.894 122.859	1.00 2.00 1.00 2.00	0
MOTA	2742	CA	VAL	285	0.979 -0.091		123.056	1.00 2.00	Ö
MOTA	2743	CB	VAL VAL	285 285	-0.952	37.714		1.00 2.00	ŏ
MOTA MOTA	2744 2745		VAL	285	0.572		123.171	1.00 2.00	Ō
MOTA	2746	C	VAL	285	0.274		122.417	1.00 2.00	0
ATOM	2747	ō	VAL	285	-0.572		121.532	1.00 2.00	0
ATOM	2748	N	ASP	286	0.609		123.026	1.00 2.00	0
MOTA	2750	CA	ASP	286	-0.056		122.663	1.00 2.00	0
MOTA	2751	CB	ASP	286	0.771		123.071	1.00 22.05 1.00 24.93	0
MOTA	2752	CG	ASP	286	1.192		124.531 125.344	1.00 24.93	ŏ
MOTA	2753		ASP	286	0.578 2.152		124.866	1.00 28.80	ŏ
MOTA	2754	OD2	ASP	286 286	-1.494	33.141		1.00 2.00	ŏ
MOTA	2755 2756	C O	ASP ASP	286	-1.993		123.881	1.00 12.72	0
MOTA MOTA	2757	N	GLU	287	-2.144	32.036	122.817	1.00 28.94	0
ATOM	2759	CA	GLU	287	-3.543	31.744		1.00 28.07	0
MOTA	2760	CB	GLU	287	-3. 91 2		122.657	1.00 59.96	0
MOTA	2761	CG	GLU	287	-3.610		121.177	1.00 66.13 1.00 70.85	0
MOTA	2762	CD	GLU	287	-2.139	29.730	119.723	1.00 70.85	Ö
MOTA	2763	OE1	GLU	287	-1.712 -1.412		121.822	1.00 73.00	Ö
ATOM	2764	OE2	GLU	287 287	-3.876		124.638	1.00 30.16	Ŏ
ATOM	2765 2766	C O	GLU	287	-5.017		125.029	1.00 62.37	0
MOTA MOTA	2767	N	THR	288	-2.852		125.455	1.00 41.83	0
MOTA	2769	CA	THR	288	-2.995		126.894	1.00 36.69	0
ATOM	2770	CB	THR	288	-2.269		127.457	1.00 11.26	0
MOTA	2771	OG1	THR	288	-0.865		127.171 126.800	1.00 11.65 1.00 11.00	0
MOTA	2773	CG2	THR	288	-2.827 -2.461		127.533	1.00 36.73	õ
MOTA	2774	C	THR THR	288 288	-2.015		128.675	1.00 18.83	0
atom atom	2775 2776	O N	LEU	289	-2.492		126.776	1.00 6.34	0
ATOM	2778	CA	LBU	289	-2.027	35.334	127.232	1.00 2.00	0
ATOM	2779	CB	LEU	289	-3.081		128.155	1.00 2.00	0
ATOM	2780	CG	LEU	289	-4.282		127.346	1.00 2.00 1.00 2.00	0
MOTA	2781		LEU	289	-5.390		128.263 126.412	1.00 2.00 1.00 2.00	Ö
MOTA	2782	_	LEU	289	-3.817 -0.615		120.412 127.826	1.00 2.00	Õ
MOTA	2783	C	LEU	289 289	-0.299	36.445	128.493	1.00 2.00	0
MOTA MOTA	2784 2785	O N	MET	290	0.244	34.475		1.00 20.85	0
ATOM	2787	CA	MET	290	1.619	34.548	128.046	1.00 23.75	0
ATOM	2788	CB	MET	290	2.238		128.212	1.00 24.81	0
ATOM	2789	CG	MET	2 9 0	3.717	33.200	128.623	1.00 23.46 1.00 26.75	Õ
MOTA .	2790	SD	MET	290	4.726	31.86/	127.935 126.310	1.00 29.16	ő
MOTA	2791	CE	MET	290	5.196 2. 43 0		127.015	1.00 19.20	0
MOTA	2792	C	MET	290 290	2.545		125.858	1.00 29.26	0
MOTA	2793 2794	0 N	MET CYS	290 291	3.003	36.454	127.449	1.00 2.00	0
ATOM ATOM	2794	CA	CYS	291	3.797	37.336	126.590	1.00 2.00	0
MOTA	2797	CB	CYS	291	3.624	38.773	127.062	1.00 17.87	0
ATOM	2798	SG	CYS	291	1.913		127.404	1.00 13.14	0
ATOM	2799	C	CYS	291	5.293		126.465	1.00 2.00	0
MOTA	2800	0	CYS	291	5.958		127.448	1.00 23.09 1.00 5.71	0
MOTA	2801	N	SER	292	5.808		125.244 124.928	1.00 9.30	0
MOTA	2803	CA	SER	292	7.223 7.385	35.709	124.928	1.00 12.57	ő
MOTA	2804	CB	SER SER	292 2 9 2	6.548		124.593	1.00 20.05	ō
MOTA	2805	OC;	JER	.,.					

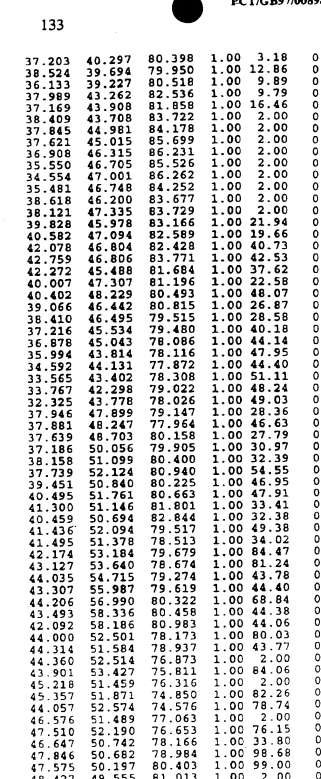
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						30 10E	124.113	1.00 4.04	0
MOTA	2807	C	SER	292	7.691 6.867	38.867	123.485	1.00 14.24	ŏ
MOTA	2808	0	SER PHE	292 293	8.993	38.487	124.128	1.00 2.00	0
MOTA	2809	N CA	PHE	293	9.526	39.623	123.385	1.00 2.00	0
MOTA	2811 2812	CB	PHE	293	10.077		124.337	1.00 17.01	0
MOTA MOTA	2813	CG	PHE	293	9.063		125.270	1.00 19.48	0
MOTA	2814	CD1		29 3	8.772		126.466	1.00 23.05 1.00 21.44	0
MOTA	2815	CD2	PHE	293	8.414		124.967 127.352	1.00 24.82	Ö
MOTA	2816	CE1	PHE	293	7.852		125.845	1.00 20.24	ŏ
MOTA	2817	CE2	PHE	293	7. 494 7. 21 3		127.044	1.00 22.28	ŏ
MOTA	2818	CZ	PHE	293 293	10.641	39.259	122.412	1.00 2.00	Ó
MOTA	2819	C O	PHE PHE	293	11.715	38.839	122.853	1.00 17.14	0
MOTA	2820 2821	N	GLN	294	10.390	39.418	121.104	1.00 2.00	0
MOTA MOTA	2823	CA	GLN	294	11.404		120.060	1.00 2.00	0
ATOM	2824	CB	GLN	294	10.748	38.742	118.743	1.00 14.58	0
MOTA	2825	CG	GLN	294	10.074		118.798	1.00 22.67 1.00 24.62	0
MOTA	2826	CD	GLN	294	8.684	37.387	118.179 118.535	1.00 27.17	ŏ
MOTA	2827		GLN	294	7.835 8.441	36.206	117.258	1.00 25.84	ŏ
MOTA	2828	NE2	GLN	294 294	12.182	40.464	119.855	1.00 2.00	Ō
MOTA	2831	C	GLN GLN	294	11.587	41.528	119.660	1.00 20.63	0
MOTA MOTA	2832 2833	O N	ILE	295	13.507	40.379	119.893	1.00 11.23	0
ATOM	2835	CA	ILE	295	14.356		119.772	1.00 12.32	0
MOTA	2836	CB	ILE	295	15.400	41.624	120.968	1.00 10.45	0
ATOM	2837	CG2	ILE	295	16.277	42.875	120.886 122.324	1.00 10.45 1.00 10.45	Ö
ATOM	2838	CG1	ILE	295	14.674 13.756	41.607 42.807	122.580	1.00 10.45	ŏ
MOTA	2839	CD1		295 295	15.126	41.703	118.453	1.00 7.79	ŏ
MOTA	2840	C	ILE	295 295	15.723	40.741	117.974	1.00 10.45	0
MOTA MOTA	2841 2842	N N	LEU	296	15.087	42.900	117.873	1.00 11.71	0
ATOM	2844	ĊA	LEU	296	15.824		116.658	1.00 13.33	0
MOTA	2845	CB	LEU	296	14.987	44.102	115.717	1.00 24.46	0
ATOM	2846	CG	LEU	296	13.835	43.467	114.946	1.00 22.89 1.00 22.12	ŏ
ATOM	2847		LEU	296	12.626		115.862 113.732	1.00 19.71	ŏ
ATOM	2848	CD2		296	13. 49 9 16. 96 5	44.070	117.224	1.00 13.82	Ō
MOTA	2849	C	LEU	296 296	16.785	45.260	117.485	1.00 13.73	0
MOTA	2850 2851	O N	LEU LYS	297	18.135	43.454	117.381	1.00 2.00	0
MOTA MOTA	2853	CA	LYS	297	19.319		117.997	1.00 2.00	0
ATOM	2854	CB	LYS	297	20.224"		118.527	1.00 79.20	0
MOTA	285 5	CG	LYS	297	20.792	42.094	117.419	1.00 79.65 1.00 76.76	Ö
ATOM	2856	CD	LYS	297	21.898	41.187 40.430	117.915 116.762	1.00 71.01	ŏ
MOTA	2857	CE	LYS	297 297	22.543 23.644	39.543	117.232	1.00 68.76	0
MOTA	2858	NZ	LYS LYS	297	20.259	45.149	117.376	1.00 2.00	0
MOTA MOTA	2862 2863	С 0	LYS	297	21.255	45.501	118.013	1.00 94.11	0
MOTA	2864	N	PRO	298	19.995	45.656	116.152	1.00 40.11	0
MOTA	2865	CD	PRO	298	18.942		115.188	1.00 2.00 1.00 40.11	0
ATOM	2866	CA	PRO	298	20.900		115.553	1.00 2.00	ŏ
MOTA	2867	CB	PRO	298	20.220 19.570	46.981	114.227 113.886	1.00 2.00	ō
MOTA	2868	CG	PRO	298 298	21,264	47.935	116.312	1.00 40.11	0
MOTA MOTA	2869 2870	C	PRO PRO	298	21.041	48.056	117.514	1.00 2.00	0
MOTA	2871	N	ALA	299	21.854	48.878	115.574	1.00 61.74	0
MOTA	2873	CA	ALA	299	22.276	50.176	116.101	1.00 61.74	0
ATOM	2874	CB	- ALA	299	23.627	50.063	116.799	1.00 2.00 1.00 61.74	Ö
MOTA	2875	С	ALA	299	22.373	51.193	114.973	1.00 61.74	Ö
MOTA	2876	0	ALA	299	21.893 41.191	52.319 29.848	115.105 91.500	1.00 48.97	ŏ
MOTA	2877	N	ASN	508 508	39.902	30.150	90.B96	1.00 48.97	Ō
MOTA	2879	CA	ASN ASN	508	38.951	30.831	91.887	1.00 0.74	0
MOTA MOTA	2880 2881	CB CG	ASN	508	37.666	31.359	91.203	1.00 0.74	0
ATOM	2882	ODI		508	36.879	30.587	90.592	1.00 0.74	0
ATOM	2883	ND2		508	37.453	32.682	91.290	1.00 0.74	0
MOTA	2886	C	ASN	508	40.096	31.041	89.687	1.00 48.97 1.00 0.74	0
MOTA	2887	0	ASN	508	40.274	32,266 30,387	89.79 8 88.533	1.00 17.27	0
ATOM	2888	N	ILE	509	40.006	/ at. vc	د د د . ن ن		~

MOTA MOTA MOTA	2890 2891 2892	CA CB CG2	ILE ILE ILE	509 509 509	40.188 40.088 38.650	30.983 29.889 29.700	87.224 86.127 85.673	1.00 18.75 1.00 44.37 1.00 52.59	0 0 0
MOTA MOTA	2893 2894	CG1	ILE	509 509	41.013 42.489	30.238 30.093	84.966 85.305	1.00 44.31 1.00 44.07	0
MOTA	2895	C	ILE	509	39.196	32.089	86.957	1.00 18.83	0
MOTA MOTA	2896 2897	O N	ILE ASP	509 510	39. 46 3 38.070	32.956 32.071	86.131 87.670	1.00 44.94 1.00 2.00	0
ATOM	2899	CA	ASP	510	37.048	33.088	87.485	1.00 2.00	0
ATOM ATOM	2900 2901	CB CG	ASP ASP	510 510	35.711 35.029	32.605 31.634	88.031 87.089	1.00 52.76 1.00 54.58	0
ATOM	2902		ASP	510	34.008	32.015	86.483	1.00 54.54	ŏ
MOTA	2903	OD2	ASP ASP	510 510	35.521 37.403	30.497 34.442	86.944 88.056	1.00 58.16 1.00 2.00	0
MOTA MOTA	2904 2905	C O	ASP	510	37.016	35.464	87.490	1.00 52.34	ŏ
MOTA	2906	N	SER	511	38.143	34.471	89.160	1.00 2.00	0
ATOM ATOM	2908 2909	CA CB	SER SER	511 511	38.548 39.372	35.763 35.570	89.739 91.015	1.00 2.00 1.00 53.95	0
ATOM	2910	OG	SER	511	38.631	34.907	92.020	1.00 50.17	0
ATOM	2912	C	SER	511 511	39. 44 7 39.228	36.379 37.488	88.690 88.203	1.00 2.00 1.00 59.62	0
ATOM ATOM	2913 2914	N N	SER	512	40.453	35.584	88.349	1.00 45.56	ŏ
MOTA	2916	CA	ILE	512	41.462	35.892	87.361	1.00 40.32	0
ATOM ATOM	2917 2918	CB CG2	ILE	512 512	42.224 43.250	34.598 34.842	87.035 85.943	1.00 2.00 1.00 2.00	0
ATOM	2919	CG1	ILE	512	42.897	34.092	88.316	1.00 2.00	0
ATOM	2920	CD1		512 512	43.635 40.843	32.772 36.514	88.174 86.105	1.00 2.00 1.00 41.22	0
MOTA MOTA	2921 2922	C O	ILE	512	41.051	37.700	85.848	1.00 2.00	ŏ
MOTA	2923	N	ILE	513	40.061	35.728	85.358	1.00 2.00 1.00 2.00	0
ATOM ATOM	2925 2926	CA CB	ILE	513 513	39.408 38.435	36.195 35.123	84.136 83.529	1.00 2.00	Ö
ATOM	2927	CG2	ILE	513	37.560	35.757	82.455	1.00 2.00	0
ATON	292B	CG1		513	39.227 38.365	33.963 32.796	82.890 . 82.367	1.00 2.00 1.00 2.00	0
ATOM ATOM	2929 2930	CD1 C	ILE	51 3 513	38.640	37.493	84.335	1.00 2.00	ŏ
MOTA	2931	ō	ILE	513	38.764	38.403	83.519	1.00 2.00	0
ATOM	2932 2934	N CA	GLN GLN	514 514	37.858 37.117	37.611 38.853	85.402 85.581	1.00 12.46 1.00 12.46	0
MOTA MOTA	2935	CB	GLN	514	35.985	38.678	86.591	1.00 26.62	Ō
MOTA	2936	CG	GLN	514	36.384	38.251 38.279	87.972 88.929	1.00 26.62 1.00 26.62	0
MOTA MOTA	2937 2938	CD OE1	GLN GLN	514 514	35.195 34.582	39.337	89.140	1.00 26.62	ŏ
ATOM	2939	NE2		514	34.851	37.116	89.502	1.00 26.62	0
ATOM	2942	C	GLN	514 514	38.009 37. - 686	40.055 41.205	85.936 85.613	1.00 12.46 1.00 26.62	0
ATOM ATOM	2943 2944	N O	GLN ARG	515	39.147	39.773	86.564	1.00 6.50	0
MOTA	2946	CA	ARG	515	40.090	40.810	86.937 87.917	1.00 7.53 1.00 8.04	0
MOTA MOTA	2947 2948	CB CG	ARG ARG	515 515	41.125 40.626	40.251 40.257	89.362	1.00 8.04	ŏ
ATOM	2949	CD	ARG	515	41.182	39.117	90.212	1.00 8.04	0
MOTA	2950	NE	ARG	51 5	42.609 43.355	39.250 38.276	90.490 90.998	1.00 8.04 1.00 8.04	0
MOTA MOTA	2952 2953	CZ NH1	ARG ARG	51 5 51 5	42.805	37.103	91.275	1.00 8.04	0
MOTA	2956	NH2	ARG	515	44.646	38.478	91.228 85.669	1.00 8.04 1.00 12.92	0
MOTA MOTA	2959 2960	0	ARG ARG	515 515	40.745 40.840	41.325 42.537	85.464	1.00 8.04	ŏ
MOTA	2961	N	LEU	516	41.167	40.398	84.810	1.00 8.93	0
ATOM	2963	CA	LEU	516 516	41.788 42.172	40.727 39.431	83.525 82.786	1.00 6.71 1.00 2.00	0
MOTA MOTA	2964 2965	CB CG	LEU LEU	516	43.298	38.561	83.385	1.00 2.00	0
MOTA	2966	CD1	LEU	516	43.057	37.096	83.083 82.843	1.00 2.00 1.00 2.00	0
MOTA MOTA	2967 2968	CD2	LEU LEU	516 516	44 .650 40.809	38.986 41.562	82.675	1.00 5.82	0
ATOM	2969	Ö	LEU	516	41.187	42.516	82.002	1.00 2.00	0
ATOM	2970	И	LEU LEU	517 517	39.53 4 38.519	41.228 41.928	82.755 81.993	1.00 2.00 1.00 2.73	0
MOTA MOTA	2972 2973	CA CB	LEU	517	37.336	40.993	81.752	1.00 7.08	Ċ

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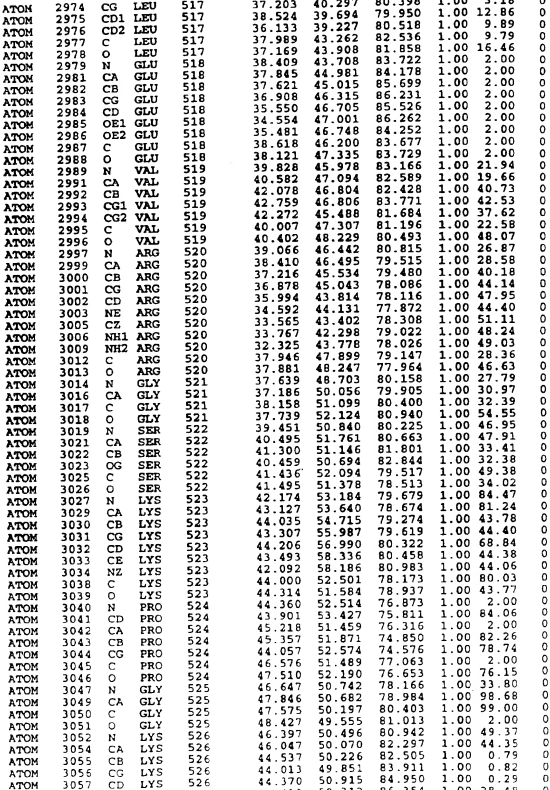
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ATOM 3064 O LYS 526 46.461 48.502 82.479 1.00 41.81 0 ATOM 3065 N ASN 527 47.372 48.337 83.412 1.00 6.60 0 ATOM 3065 N ASN 527 47.861 46.972 83.653 1.00 6.60 0 ATOM 3067 CA ASN 527 47.861 46.972 83.653 1.00 6.60 0 ATOM 3068 CB ASN 527 50.300 47.733 83.645 1.00 30.42 0 ATOM 3068 CB ASN 527 50.300 47.733 83.655 1.00 30.42 0 ATOM 3070 0D1 ASN 527 50.300 47.733 83.655 1.00 30.42 0 ATOM 3070 0D1 ASN 527 50.300 47.733 83.655 1.00 30.42 0 ATOM 3070 0D1 ASN 527 50.300 47.733 83.665 1.00 30.42 0 ATOM 3070 ASN 527 50.300 47.733 83.665 1.00 30.42 0 ATOM 3070 ASN 527 50.500 47.733 83.665 1.00 30.42 0 ATOM 3076 N ASN 527 46.895 46.325 84.463 1.00 5.00 17 0 ATOM 3076 N AVAL 528 46.541 43.874 84.601 1.00 26.67 0 ATOM 3078 CA VAL 528 46.541 43.874 85.443 1.00 26.67 0 ATOM 3080 CG VAL 528 46.541 43.874 85.443 1.00 26.67 0 ATOM 3080 CG VAL 528 44.527 43.866 83.956 1.00 33.41 0 ATOM 3081 CG2 VAL 528 44.527 43.858 83.956 1.00 33.61 0 ATOM 3081 CG2 VAL 528 44.524 41.856 83.956 1.00 35.61 0 ATOM 3082 C VAL 528 44.527 42.400 86.018 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.547 42.940 86.018 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.547 42.940 86.018 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.604 42.207 88.956 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.604 42.207 88.956 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.604 42.207 88.956 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.228 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.228 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.228 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.228 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.228 1.00 35.61 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.238 1.00 35.00 0 ATOM 3080 CG GL VAL 528 40.08 43.237 87.238 1.00 32.00 0 ATOM 3080 CG GL VAL 528 48.08 48.08 48.09 1.00 35.61 0 ATOM 3080 CG GL VAL 528 48.08 48.08 1.00 32.29 0 ATOM 3080 CG GL VAL 528 48.08 48.08 1.00 32.29 0 ATOM 3080 CG GL VAL 528 48.08 48.08 1.00 32.29 0 ATOM 3080 CG GL VAL 528 48.08 48.08 1.00 32.90 0 ATOM 3090 CG EGL 528 50.00 0 ATOM 3090 CG EGL 528 50.00 0 ATOM 3090	ATOM	3059	NZ	LYS	526	45.166	51.143	87.348	1.00 28.16	0
ATOM 3065 N ASN 527 47.372 48.337 83.412 1.00 6.60 0 ATOM 3068 CB ASN 527 47.861 46.977 84.381 1.00 20.38 ATOM 3069 CG ASN 527 50.300 47.733 83.645 1.00 30.42 0 ATOM 3070 0D1 ASN 527 50.695 48.820 84.068 1.00 29.99 ATOM 3071 ND2 ASN 527 50.695 84.820 84.068 1.00 29.99 ATOM 3071 ND2 ASN 527 50.695 84.820 84.068 1.00 29.99 ATOM 3071 ND2 ASN 527 50.695 84.820 84.068 1.00 29.99 ATOM 3074 C ASN 527 50.695 84.820 84.068 1.00 29.99 ATOM 3074 C ASN 527 50.695 84.820 84.068 1.00 29.99 ATOM 3075 O ASN 527 45.8666 46.628 84.924 1.00 23.47 0 ATOM 3075 O ASN 527 45.8666 46.628 84.924 1.00 23.47 0 ATOM 3076 N VAL 528 46.541 43.874 85.443 1.00 26.67 0 ATOM 3078 CA VAL 528 46.541 43.874 85.443 1.00 26.67 0 ATOM 3078 CA VAL 528 44.999 41.956 85.488 1.00 33.01 0 ATOM 3080 CG1 VAL 528 44.999 41.956 85.488 1.00 33.01 0 ATOM 3083 O VAL 528 47.677 42.940 85.348 1.00 35.61 0 ATOM 3083 O VAL 528 47.677 42.940 85.348 1.00 36.65 0 ATOM 3083 O VAL 528 49.064 43.203 85.388 1.00 35.61 0 ATOM 3080 CG GGN 529 49.664 12.207 87.328 1.00 2.00 0 ATOM 3080 CG GGN 529 49.664 13.207 87.328 1.00 2.00 0 ATOM 3080 CG GGN 529 49.664 13.207 87.328 1.00 30.41 0 ATOM 3080 CG GGN 529 52.380 42.558 88.558 1.00 30.41 0 ATOM 3080 CG GGN 529 52.380 42.558 88.558 1.00 30.41 0 ATOM 3080 CG GGN 529 52.259 48.364 43.277 88.797 1.00 2.00 0 ATOM 3080 CG GGN 529 52.259 48.364 30.208 89.942 1.00 2.00 0 ATOM 3091 REZ GGN 529 52.380 42.558 88.558 1.00 30.17 0 ATOM 3098 CA LEU 530 47.545 41.689 89.663 1.00 30.49 0 ATOM 3099 CD GGN 529 52.380 42.558 88.558 1.00 30.19 0 ATOM 3099 CD GGN 529 52.380 42.558 88.558 1.00 30.19 0 ATOM 3090 CG LEU 530 48.664 37.943 89.002 88.303 1.00 2.00 0 ATOM 3090 CG LEU 530 48.664 39.002 88.303 1.00 2.00 0 ATOM 3090 CG LEU 530 48.664 39.002 88.303 1.00 2.00 0 ATOM 3090 CG LEU 530 49.665 30.90 30.90 0 ATOM 3090 CG LEU 530 49.665 30.90 30.90 0 ATOM 3090 CG LEU 530 49.665 30.90 90.00 1.00 2.00 0 ATOM 3090 CG LEU 530 49.665 30.90 90.00 1.00 2.00 0 ATOM 3090 CG LEU 530 49.665 30.90 90.00 1.00 2.00 0 ATOM 3090 CG LEU 530 49.665 30.90 90.00 1.00 2.0					526	46.461	48.602	82.479		-
ATOM 3068 CB ASN 527			-							
ATOM 3068 CE ASN 527			-							
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TOM 3076 **N **VAL **528 **47.271 **44.866 **84.647 **1.00 **23.47 **O **ATOM** 3076 **N **VAL **528 **47.271 **44.866 **84.647 **1.00 **26.67 **O **ATOM** 3079 **CB **VAL **528 **45.575 **43.004 **84.603 **1.00 **33.01 **CG **VAL **528 **45.575 **43.004 **84.603 **1.00 **33.01 **CG **VAL **528 **44.599 **41.956 **85.488 **1.00 **33.01 **CG **VAL **528 **44.599 **41.956 **85.488 **1.00 **33.01 **CG **VAL **528 **44.524 **43.858 **83.956 **1.00 **35.61 **CG **ATOM** 3083 **O **VAL **528 **48.054 **2.003 **85.348 **1.00 **36.85 **O **ATOM** 3083 **O **VAL **528 **48.054 **2.003 **85.348 **1.00 **36.85 **O **ATOM** 3086 **CG **CIN **529 **49.061 **42.407 **87.929 **1.00 **2.00 **O **ATOM** 3089 **CG **CIN **529 **49.968 **43.297 **88.787 **1.00 **30.41 **O **ATOM** 3089 **CG **CIN **529 **51.142 **42.569 **89.428 **1.00 **30.31 **O **ATOM** 3089 **CD **GIN **529 **52.380 **42.558 **88.558 **1.00 **31.37 **O **ATOM** 3099 **CD **GIN **529 **52.380 **42.558 **89.563 **1.00 **31.37 **O **ATOM** 3091 **NE2 **GIN **529 **47.545 **41.689 **89.663 **1.00 **32.34 **O **ATOM** 3095 **O **GIN **529 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 31095 **O **GIN **529 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 3109 **CD **LEU **530 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 3109 **CD **LEU **530 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 3100 **CG **LEU **530 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 3100 **CG **LEU **530 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 3100 **CG **LEU **530 **47.545 **41.689 **89.663 **1.00 **32.29 **O **ATOM** 3100 **CG **LEU **530 **47.545 **41.689 **89.663 **1.00 **22.00 **O **ATOM** 3100 **CG **LEU **530 **47.545 **41.689 **89.663 **1.00 **22.00 **O **ATOM** 3100 **CG **LEU **530 **47.545 **41.689 **89.663 **1.00 **22.00 **O **ATOM*** 3100 **CG **L										_
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ATOM 3120 CB GLU 532 54.056 35.623 91.336 1.00 19.01 0 ATOM 3121 CG GLU 532 55.176 34.623 90.966 1.00 22.74 0 ATOM 3122 CD GLU 532 56.496 34.871 91.746 1.00 25.50 0 ATOM 3123 OE1 GLU 532 56.845 36.042 92.058 1.00 20.16 0 ATOM 3124 OE2 GLU 532 57.195 33.879 92.051 1.00 20.82 0 ATOM 3125 C GLU 532 52.345 33.866 90.900 1.00 2.00 0 ATOM 3126 O GLU 532 52.833 33.059 90.116 1.00 17.28 0 ATOM 3127 N ASN 533 51.568 33.509 91.918 1.00 26.78 0 ATOM 3129 CA ASN 533 51.181 32.120 92.127 1.00 27.69 0 ATOM 3130 CB ASN 533 50.751 31.857 93.587 1.00 42.65 0 ATOM 3131 CG ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3132 OD1 ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.06 0 ATOM 3136 C ASN 533 48.568 32.605 94.387 1.00 50.92 0 ATOM 3137 O ASN 533 48.568 32.605 94.387 1.00 50.92 0 ATOM 3138 N GLU 534 48.568 32.777 90.815 1.00 24.76 0 ATOM 3138 N GLU 534 49.918 30.615 90.747 1.00 41.80 0 ATOM 3134 CB GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3141 CB GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.066 0 ATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.066 0										
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ATOM 3122 CD GLU 532 56.496 34.871 91.746 1.00 20.16 0 ATOM 3123 OE1 GLU 532 56.845 36.042 92.058 1.00 20.16 0 ATOM 3124 OE2 GLU 532 57.195 33.879 92.051 1.00 20.82 0 ATOM 3125 C GLU 532 52.345 33.866 90.900 1.00 2.00 0 ATOM 3126 O GLU 532 52.833 33.059 90.116 1.00 17.28 0 ATOM 3127 N ASN 533 51.568 33.509 91.918 1.00 26.78 0 ATOM 3129 CA ASN 533 51.181 32.120 92.127 1.00 27.69 0 ATOM 3130 CB ASN 533 50.751 31.857 93.587 1.00 42.65 0 ATOM 3131 CG ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3132 OD1 ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.06 0 ATOM 3136 C ASN 533 48.568 32.605 94.387 1.00 50.92 0 ATOM 3137 O ASN 533 49.918 30.615 90.747 1.00 41.80 0 ATOM 3138 N GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3138 N GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3142 CG GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3144 OE1 GLU 534 46.909 34.997 91.890 1.00 23.75 0 ATOM 3144 OE1 GLU 534 46.909 34.997 91.890 1.00 23.75 0				-			34.623			
ATOM 3123 OE1 GLU 532 57.195 33.879 92.051 1.00 20.82 0 ATOM 3125 C GLU 532 52.345 33.866 90.900 1.00 2.00 ATOM 3126 O GLU 532 52.833 33.059 90.116 1.00 17.28 OATOM 3127 N ASN 533 51.568 33.509 91.918 1.00 26.78 OATOM 3129 CA ASN 533 51.568 33.509 91.918 1.00 26.78 OATOM 3130 CB ASN 533 50.751 31.857 93.587 1.00 42.65 OATOM 3131 CG ASN 533 49.834 32.944 94.154 1.00 49.17 OATOM 3132 OD1 ASN 533 49.834 32.944 94.154 1.00 49.17 OATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.06 OATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.06 OATOM 3136 C ASN 533 48.568 32.605 94.387 1.00 26.24 OATOM 3137 OASN 533 49.918 30.615 90.747 1.00 41.80 OATOM 3138 N GLU 534 49.918 30.615 90.747 1.00 41.80 OATOM 3138 N GLU 534 48.146 32.592 89.893 1.00 19.83 OATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 19.83 OATOM 3141 CB GLU 534 46.509 33.924 91.239 1.00 23.75 OATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 23.75 OATOM 3144 OE1 GLU 534 46.509 33.924 91.239 1.00 23.75 OATOM 3144 OE1 GLU 534 46.509 33.924 91.239 1.00 23.75 OATOM 3144 OE1 GLU 534 46.509 33.924 91.239 1.00 23.75 OATOM 3144 OE1 GLU 534 46.990 34.997 91.890 1.00 24.06 OATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.06 OATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.06		3122	CD							
ATOM 3125 C GLU 532 52.345 33.866 90.900 1.00 2.00 0 ATOM 3126 O GLU 532 52.833 33.059 90.116 1.00 17.28 0 ATOM 3127 N ASN 533 51.568 33.509 91.918 1.00 26.78 0 ATOM 3129 CA ASN 533 51.181 32.120 92.127 1.00 27.69 0 ATOM 3130 CB ASN 533 50.751 31.857 93.587 1.00 42.65 0 ATOM 3131 CG ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3132 OD1 ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.06 0 ATOM 3136 C ASN 533 48.568 32.605 94.387 1.00 50.92 0 ATOM 3137 O ASN 533 48.568 32.605 94.387 1.00 26.24 0 ATOM 3138 N GLU 534 48.568 32.605 90.747 1.00 41.80 0 ATOM 3138 N GLU 534 49.253 32.777 90.815 1.00 24.76 0 ATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3144 OEI GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.490 34.997 91.890 1.00 24.06 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 24.06 0										_
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ATOM 3129 CA ASN 533 51.181 32.120 92.127 1.00 27.69 0 ATOM 3130 CB ASN 533 50.751 31.857 93.587 1.00 42.65 0 ATOM 3131 CG ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3132 OD1 ASN 533 50.274 34.070 94.392 1.00 50.06 0 ATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.92 0 ATOM 3136 C ASN 533 48.568 32.605 94.387 1.00 26.24 0 ATOM 3137 O ASN 533 49.918 30.615 90.747 1.00 41.80 0 ATOM 3138 N GLU 534 49.918 30.615 90.747 1.00 41.80 0 ATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 47.228 33.801 89.918 1.00 17.15 0 ATOM 3142 CG GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 25.22 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 24.06 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 29.17 0				GLU	532				1.00 17.28	
ATOM 3130 CB ASN 533									1.00 27.69	0
ATOM 3131 CG ASN 533 49.834 32.944 94.154 1.00 49.17 0 ATOM 3132 OD1 ASN 533 50.274 34.070 94.392 1.00 50.06 0 ATOM 3133 ND2 ASN 533 48.568 32.605 94.387 1.00 50.92 0 ATOM 3136 C ASN 533 50.053 31.770 91.165 1.00 26.24 0 ATOM 3137 O ASN 533 49.918 30.615 90.747 1.00 41.80 0 ATOM 3138 N GLU 534 49.918 30.615 90.747 1.00 41.80 0 ATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 47.228 33.801 89.918 1.00 17.15 0 ATOM 3142 CG GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3144 OEI GLU 534 46.400 34.997 91.890 1.00 24.06 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 29.17 0									1.00 42.65	
ATOM 3132 OD1 ASN 533						49.83	4 32.944		1.00 49.17	
ATOM 3136 C ASN 533 50.053 31.770 91.165 1.00 26.24 0 ATOM 3136 C ASN 533 50.053 31.770 91.165 1.00 24.76 0 ATOM 3137 O ASN 533 49.918 30.615 90.747 1.00 41.80 0 ATOM 3138 N GLU 534 49.253 32.777 90.815 1.00 24.76 0 ATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 47.228 33.801 89.918 1.00 17.15 0 ATOM 3142 CG GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3144 OEI GLU 534 45.598 35.132 91.323 1.00 25.22 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 24.06 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 29.17 0	MOTA	3132								
ATOM 3136 C ASN 533 49.918 30.615 90.747 1.00 41.80 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1									1.00 26.24	0
ATOM 3138 N GLU 534 49.253 32.777 90.815 1.00 24.76 0 ATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 19.83 0 ATOM 3141 CB GLU 534 47.228 33.801 89.918 1.00 17.15 0 ATOM 3142 CG GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 45.598 35.132 91.323 1.00 25.22 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 24.06 0 ATOM 3144 OEI GLU 534 44.490 36.215 90.837 1.00 29.17							8 30.615	90.747		
ATOM 3140 CA GLU 534 48.146 32.592 89.893 1.00 17.15 0 ATOM 3141 CB GLU 534 47.228 33.801 89.918 1.00 17.15 0 ATOM 3142 CG GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3143 CD GLU 534 45.598 35.132 91.323 1.00 25.22 0 ATOM 3144 OEI GLU 534 44.490 34.997 91.890 1.00 24.06 0 ATOM 3144 OEI GLU 534 44.490 36.215 90.837 1.00 29.17 0					534	49.25	3 32.777			
ATOM 3141 CB GLU 534 46.509 33.924 91.239 1.00 23.75 0 ATOM 3142 CG GLU 534 45.598 35.132 91.323 1.00 25.22 0 ATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.06 0 ATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 29.17 0	MOTA	3140	CA	GLU					1.00 17.15	
ATOM 3143 CD GLU 534 45.598 35.132 91.323 1.00 25.22 0 ATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.06 0						47.22			1.00 23.75	0
ATOM 3144 OE1 GLU 534 44.490 34.997 91.890 1.00 24.00 0							8 35.132		1.00 25.22	
				GLU	534				1.00 24.06	
	MOTA	3145	OE2	g GLU	534	45.99	1 30.21	, , , ,	.,	-

		_	CT 11	534	48.712	32,365	88.510	1.00 21.16	0
MOTA	3146 3147	С 0	GLU GLU	534	48.235	31.500	87.779	1.00 14.12	ō
ATOM ATOM	3148	N	ILE	535	49.747	33.122	88.156	1.00 2.00	0
MOTA	3150	CA	ILE	535	50.389	32.942	86.859	1.00 2.00	0
ATOM	3151	CB	ILE	535	51.442	34.023	86.570	1.00 2.00	0
MOTA	3152		ILE	535	52.050	33.784	85.190	1.00 2.00 1.00 2.00	0
MOTA	3153	CG1	ILE	535	50.793	35.411	86.641 85.849	1.00 2.00	0
MOTA	3154		ILE	535 535	51.537 51.060	36.488 31.565	86.797	1.00 2.00	ő
MOTA	3155	Ç	ILE	535	50.788	30.780	85.876	1.00 2.00	Ö
ATOM	3156 3157	O N	ARG	536	51.914	31.256	87.778	1.00 2.00	ō
MOTA	3159	CA	ARG	536	52.583	29.951	87.793	1.00 2.00	0
ATOM ATOM	3160	CB	ARG	536	53.495	29.818	89.011	1.00 39.13	0
ATOM	3161	CG	ARG	536	52.808	29.604	90.341	1.00 45.43	0
ATOM	3162	CD	ARG	536	53.839	29.648	91.474	1.00 49.26	0
MOTA	3163	NE	ARG	536	55.059	28.910	91.132	1.00 56.04	0
ATOM	3165	CZ	ARG	536	55.110	27.603	90.876 90.920	1.00 55.24 1.00 53.92	0
MOTA	3166		ARG	536	54.011 56.267	26.860 27.032	90.576	1.00 44.45	ŏ
MOTA	3169	NH2	ARG	536 536	51.562	28.811	87.742	1.00 2.00	ŏ
ATOM	3172 3173	C O	ARG ARG	536	51.806	27.778	87.126	1.00 37.01	0
ATOM ATOM	3174	N	GLY	537	50.406	29.033	88.363	1.00 2.00	0
ATOM	3176	CA	GLY	537	49.345	28.046	88.340	1.00 2.00	٥
ATOM	3177	Č.	GLY	537	48.690	28.000	86.970	1.00 2.00	0
ATOM	3178	0	GLY	537	48.292	26.924	86.512	1.00 2.00	0
MOTA	3179	N	LEU	538	48.574	29.164	86.319	1.00 9.51 1.00 7.63	0
MOTA	3181	CA	LEU	538	47.979	29.276 30.736	84.977 84.513	1.00 7.63 1.00 2.00	ő
ATOM	3182	CB	LEU	538 538	47.930 46.654	31.524	84.772	1.00 2.00	ŏ
MOTA	3183	CG	Leu Leu	538	46.865	32.966	84.408	1.00 2.00	ŏ
MOTA	3184 3185		LEU	538	45.531	30.940	83.972	1.00 2.00	0
MOTA MOTA	3186	C	LEU	538	48.816	28.472	84.000	1.00 12.73	0
ATOM	3187	ŏ	LEU	538	48.295	27.635	83.271	1.00 2.00	0
ATOM	3188	N	CYS	539	50.120	28.724	84.000	1.00 2.00	0
MOTA	3190	CA	CYS	539	51.036	28.001	83.132	1.00 2.00 1.00 20.52	0
MOTA	3191	CB	CYS	539	52.473	28.494	83.337 83.567	1.00 20.32	ő
ATOM	3192	SG	CYS	539 539	52.713 50.957	30.280 26.501	83.474	1.00 2.00	ŏ
MOTA	3193	C O	CYS CYS	53 <i>9</i> 539	50.854	25.658	82.588	1.00 29.69	Ō
MOTA MOTA	3194 3195	N	LEU	540	50.984	26.191	84.769	1.00 26.60	0
MOTA	3197	CA	LEU	540	50.919	24.819	85.265	1.00 26.60	0
MOTA	3198	CB	LBU	540	51.106	24.818	86.786	1.00 2.00	0
ATOM	3199	CG	LEU	540	52.539	24.770	87.339	1.00 2.00	0
ATOM	3200		LEU	540	53.001	23.360	87.385	1.00 2.00 1.00 2.00	Ö
MOTA	3201		LEU	540	53.492	25.577 24.061	86.497 84.899	1.00 26.60	ő
MOTA	3202	C	LEU	540	49.635 49.677	22.856	84.619	1.00 2.00	ŏ
MOTA	3203	N O	Leu Lys	540 541	48.500	24.756	84.901	1.00 13.81	0
MOTA MOTA	3204 3206	CA	LYS	541	47.219	24.126	84.561	1.00 17.46	0
MOTA	3207	CB	LYS	541	46.046	24.964	85.108	1.00 14.61	0
MOTA	3208	CG	LYS	541	45.844	24.880	86.624	1.00 22.53	0
MOTA	3209	CD	LYS	541	44.709	25.777	87.150	1.00 31.64 1.00 37.15	0 0
ATOM	3210	CE	LYS	541	45.175	27.201	87.517 88.284	1.00 37.13	ŏ
MOTA	3211	NZ	LYS	541	44.147 47.047	28.010 23.891	83.046	1.00 16.97	ō
MOTA	3215	C	LYS	541 541	46.862	22.745	82.608	1.00 8.92	0
MOTA MOTA	3216 3217	О И	LYS S E R	542	47.131	24.963	82.253	1.00 2.00	0
ATOM	3217	CA	SER	542	46.975	24.877	80.791	1.00 2.00	0
ATOM	3220	CB	SER	542	47.165	26.256	80.150	1.00 2.00	0
ATOM	3221	0G	SER	542	48.499	26.703	80.298	1.00 2.00	0
ATOM	3223	С	SER	542	47.915	23.870	80.107	1.00 2.00	0
MOTA	3224	0	SER	542	47.450	22.991	79.377	1.00 2.00 1.00 8.64	0
MOTA	3225	N	ARG	543	49.223 50.244	24.016 23.128	80.338 79.772	1.00 8.64 1.00 8.64	0
ATOM	3227	CA	ARG	543 543	51.607	23.128	80.434	1.00 2.00	Ö
MOTA	3228 3229	CB CG	ARG ARG	543	52. 67 6	22.339	80.116	1.00 4.46	ō
MOTA MOTA	3229	CD	ARG	543	53. 7 57	22.313	81.173	1.00 2.00	0
ATOM	3231	NE	ARG	543	54.989	21.663	80.708	1.00 2.00	O

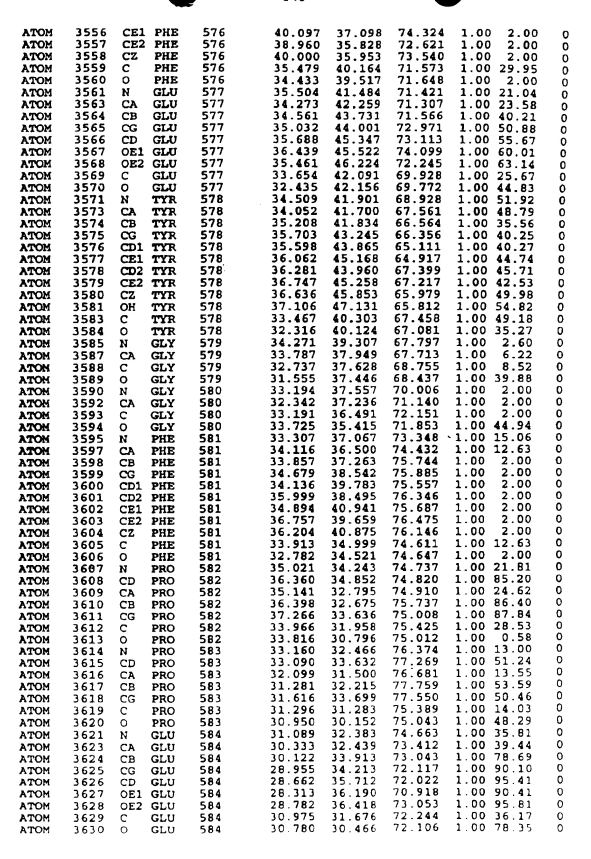
MOTA MOTA	3233 3234	CZ NH1	ARG ARG	543 543	55.326 54.523	20.395 19.606	80.952 81.656	1.00 2.00 1.00 3.99	0
ATOM	3237	NH2	ARG	543	56.483	19.915	80.508	1.00 2.00	Ō
ATOM	3240	С	ARG	543	49.873	21.660	79.981	1.00 8.64	0
ATOM	3241	0	ARG	543	50.187	20.806	79.148	1.00 5.52	0
MOTA	3242	N	GLU	544	49.227	21.352	81.098	1.00 42.94	0
ATOM	3244	CA	GLU	544	48.834	19.978	81.355	1.00 44.22	0
MOTA	3245	CB	GLU	544	48.308	19.841	82.778	1.00 37.01	0
MOTA	3246	CG	GLU	544	48.175	18.415	83.249	1.00 41.48	0
MOTA	3247	CD	GLU	544	47.561	18.341 18.858	84.626 84.791	1.00 46.37 1.00 53.09	0
MOTA	3248		GLU	5 44 5 44	46.433 48.202	17.777	85.543	1.00 47.19	Ö
MOTA	3249		GLU	544	47.763	19.584	80.339	1.00 40.49	Ö
MOTA	3250 3251	C	GLU	544	47.742	18.450	79.867	1.00 34.80	ŏ
ATOM ATOM	3252	N	ILE	545	46.898	20.538	79.994	1.00 2.00	Ŏ
ATOM	3254	CA	ILE	545	45.823	20.315	79.017	1.00 2.00	0
ATON	3255	CB	ILE	545	44.805	21.487	78.959	1.00 11.87	0
ATOM	3256	CG2	ILE	545	43.645	21.102	78.064	1.00 6.53	0
MOTA	3257	CG1	ILE	545	44.288	21.836	80.353	1.00 16.65	0
MOTA	3258	CD1		545	43.267	22.955	80.361	1.00 13.01	0
MOTA	3259	C	ILE	545	46.377	20.149	77.605	1.00 2.00	0
MOTA	3260	0	ILE	545	45.960	19.267 21.018	76.868 77.227	1.00 9.05 1.00 2.00	0
MOTA	3261	N	PHE	546	47.302 47.895	20.945	75.918	1.00 2.00	Ö
ATOM	3263	CA	PHE	546 546	49.058	21.906	75.816	1.00 2.00	ŏ
MOTA	3264 3265	CB CG	PHE PHE	546	48.653	23.335	75.875	1.00 2.00	ŏ
ATOM ATOM	3266		PHE	546	49.512	24.293	76.399	1.00 2.00	Ō
ATOM	3267		PHE	546	47.411	23.730	75.426	1.00 2.00	0
ATOM	3268			546	49.127	25.630	76.473	1.00 2.00	0
ATOM	3269		PHE	546	47.023	25.050	75.496	1.00 2.00	0
MOTA	3270	\mathbf{cz}	PHE	546	47.881	26.006	76.020	1.00 2.00	0
ATOM	3271	С	PHE	546	48.374	19.538	75.658	1.00 2.00 1.00 2.00	0
MOTA	3272	0	PHE	546	48.141	18.990	74.596 76.647	1.00 2.00 1.00 2.00	ŏ
MOTA	3273	N	LEU	547	49.012 49.527	18.935 17.582	76.506	1.00 2.00	ŏ
ATOM	3275	CA	LEU	547 547	50.499	17.289	77.654	1.00 6.91	ŏ
MOTA MOTA	3276 3277	CB CG	LEU	547	51.754	18.159	77.582	1.00 8.40	
ATOM	3411						11.302	1.00 0.40	0
	マクフR			-		18.701	78.934	1.00 16.17	ŏ
	3278 3279	CD1	LEU	547 547	52.096 52.905	18.701 17.363	78.934 77.029	1.00 16.17 1.00 11.72	0
MOTA MOTA	3278 3279 3280	CD1		547 547 547	52.096 52.905 48. 42 2	18.701 17.363 16.526	78.934 77.029 76.434	1.00 16.17 1.00 11.72 1.00 2.00	0 0 0
MOTA	3279	CD1 CD2	TEA TEA TEA	547 547 547 547	52.096 52.905 48.422 48.642	18.701 17.363 16.526 15.414	78.934 77.029 76.434 75.946	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91	0 0 0
MOTA MOTA	3279 3280 3281 3282	CD1 CD2 C O N	LEU LEU LEU SER	547 547 547 547 548	52.096 52.905 48.422 48.642 47.230	18.701 17.363 16.526 15.414 16.881	78.934 77.029 76.434 75.946 76.903	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34	0 0 0 0
MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284	CD1 CD2 C O N CA	LEU LEU LEU SER SER	547 547 547 547 548 548	52.096 52.905 48.422 48.642 47.230 46.091	18.701 17.363 16.526 15.414 16.881 15.964	78.934 77.029 76.434 75.946 76.903 76.898	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 5.34	0 0 0 0 0
MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285	CD1 CD2 C O N CA CB	LEU LEU LEU SER SER SER	547 547 547 547 548 548 548	52.096 52.905 48.422 48.642 47.230 46.091 45.099	18.701 17.363 16.526 15.414 16.881 15.964 16.356	78.934 77.029 76.434 75.946 76.903 76.898 77.993	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 5.34 1.00 36.19	0 0 0 0 0 0
ATOM MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286	CD1 CD2 C O N CA CB OG	LEU LEU LEU SER SER SER SER	547 547 547 547 548 548 548 548	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64	0 0 0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286 3288	CD1 CD2 C O N CA CB OG C	LEU LEU LEU SER SER SER SER SER	547 547 547 547 548 548 548 548 548	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975	78.934 77.029 76.434 75.946 76.903 76.898 77.993	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 5.34 1.00 36.19 1.00 39.64	000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286 3288 3289	CD1 CD2 C O N CA CB OG C	LEU LEU LEU SER SER SER SER SER	547 547 547 547 548 548 548 548 548	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374 44.299	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 5.34	0000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286 3288 3289 3290	CD1 CD2 C O N CA CB OG C	LEU LEU LEU SER SER SER SER SER SER	547 547 547 547 548 548 548 548 548	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 5.34 1.00 5.34 1.00 23.79 1.00 23.79	00000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286 3288 3289 3290 3292	CD1 CD2 C O N CA CB OG C O N CA	LEU LEU LEU SER SER SER SER SER SER GLN GLN	547 547 547 547 548 548 548 548 548 548 549	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.384 44.299 45.384	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 23.94 1.00 46.80	000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286 3288 3289 3290	CD1 CD2 C O N CA CB OG C	LEU LEU LEU SER SER SER SER SER SER	547 547 547 548 548 548 548 548 549 549 549	52.096 52.905 48.422 48.642 47.230 45.099 45.611 45.374 44.299 45.968 45.368	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.354	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 246.80 1.00 38.43	0000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3286 3288 3289 3299 32992 32993 3294 3295	CD1 CD2 C O N CA CB OG C O N CA CB CCB CCD	LEU LEU LEU SER SER SER SER SER GLI GLI GLI GLI	547 547 547 548 548 548 548 549 549 549	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374 44.299 45.968 45.384 44.751 43.567 43.180	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.94 1.00 46.80 1.00 38.43 1.00 38.53	0 0 0 0 0 0 0 0 0 0 0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3288 3288 3288 3289 3290 3299 32993 32994 3295 3296	CD1 CD2 C O N CA CB OG C O N CA CB CD OE1	LEU LEU LEU SER SER SER SER GLIN GLIN GLIN GLIN GLIN	547 547 547 548 548 548 548 549 549 549 549	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374 44.299 45.384 44.751 43.567 43.180 43.934	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.624	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.94 1.00 46.80 1.00 38.43 1.00 38.53 1.00 46.42	0 0 0 0 0 0 0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3288 3289 3290 3299 32993 32995 3297	CD1 CD2 C O N CA CB OG C O N CA CB CG CD OE1 NE2	LEU LEU LEU SER SER SER GLIIN GLIIN GCLIN GCLIN GCLIN	547 547 547 548 548 548 548 549 549 549 549 549	52.096 52.905 48.422 48.642 47.230 46.091 45.611 45.374 44.299 45.968 45.384 44.7567 43.567 43.180 43.934 42.002	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.595	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 46.80 1.00 38.43 1.00 38.53 1.00 38.53 1.00 39.54	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3288 3288 3289 3290 3299 32993 32995 3297 3300	CD1 CD2 C O N CA CB OG C O N CA CB CG CD OE1 NE2 C	LEU LEU LEU SER SER SER GLIN GCLIN GCLIN GCLIN GCLIN	547 547 547 548 548 548 549 549 549 549 549 549	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374 44.299 45.968 45.384 44.7567 43.180 43.934 42.002 46.456	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074 16.547	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.624 74.595 72.182	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.94 1.00 46.80 1.00 38.43 1.00 38.53 1.00 46.42	0 0 0 0 0 0 0 0 0 0 0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3288 3288 3289 3299 3299 32995 32995 3297 3300 3301	CD1 CD2 C O N CA CB OG C O N CA CB CG CD OE1 N CA CB CG CO O O	LEU LEU LEU SER SER SER GLIN GCLIN GCLIN GCLIN GCLIN GCLIN GCLIN	547 547 547 548 548 548 549 549 549 549 549 549 549	52.096 52.905 48.422 48.642 47.230 46.091 45.611 45.374 44.299 45.968 45.384 44.751 43.567 43.180 43.934 42.002 46.456 47.634	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.672 16.547 16.766	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.624 74.595 72.182 72.451	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 36.66 1.00 23.79 1.00 23.79 1.00 46.80 1.00 38.43 1.00 38.53 1.00 38.53 1.00 39.54 1.00 39.54 1.00 25.87	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3284 3285 3288 3299 3299 3299 32995 32997 3300 3301 3302	CD1 CD2 CONCACBOC CONCACBCCCONE2 CONCACBCCCONE2	LEU LEU LEU SER SER SER SER GLIN GLIN GLIN GLIN GLIN GLIN PRO	547 547 547 548 548 548 549 549 549 549 5549 5549 5	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.968 45.384 44.751 43.567 43.180 43.934 42.002 46.456 47.634 46.068	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074 16.547	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.624 74.595 72.182	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 46.80 1.00 38.43 1.00 38.53 1.00 46.42 1.00 39.54 1.00 39.54 1.00 25.87 1.00 201 1.00 2.00 1.00 36.68	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3288 3288 3288 3299 3299 3299 3299	CD1 CD2 CONCACBOC CONCACCONCCONCCONCCONCC	LEU LEU LEU SER SER SER SER GLIN GLIN GLIN PRO PRO	547 547 547 548 548 548 548 549 5549 5549 5549 555 555 555 555 555	52.096 52.905 48.422 48.642 47.230 46.091 45.611 45.374 44.299 45.968 45.384 44.751 43.567 43.180 43.934 42.002 46.456 47.634	18.701 17.363 16.526 15.414 16.881 15.964 17.389 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074 16.766 16.766 16.105	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.990 74.054 73.624 74.595 72.182 72.451 70.963 70.469 69.921	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 23.94 1.00 38.43 1.00 38.53 1.00 46.42 1.00 39.54 1.00 25.87 1.00 42.01 1.00 2.00 1.00 36.68 1.00 2.00	000000000000000000000000000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	3279 3280 3281 3282 3288 3288 3288 3299 32992 32993 32995 32996 32997 33001 33002 3303 3304	CD1 CD2 C O N CAB CG CD OE1 C O N CAB CG CD OE2 C O N CAB CG CD OE2	LEU LEU LEU SER SER SER GLIN GLIN GRO GRO PRO	547 547 547 548 548 548 549 549 549 549 5549 5549 5	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.384 44.751 43.567 43.180 43.934 42.002 46.456 47.634 46.068 47.075 46.253	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074 16.547 16.766 16.105 15.738 15.738 15.738	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.090 74.054 73.624 74.595 72.182 72.451 70.963 70.963 70.469 69.921 68.750	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 23.94 1.00 38.43 1.00 38.53 1.00 46.80 1.00 39.54 1.00 39.54 1.00 25.87 1.00 42.01 1.00 2.00 1.00 36.68 1.00 36.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3288 3288 3288 3289 3299 3299 3299 3299	CD1 CD2 CONCACGCCD OE1 CCACCGCCD OE2 CCACCB	LEU LEU LEU SER SER SER SER GLIN GLIN GLIN PRO PRO	547 547 547 548 548 548 548 549 5549 5549 555 555 555 555 555 555	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.384 44.299 45.384 44.567 43.180 43.934 44.729 46.456 47.634 46.729 47.075 44.886	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074 16.547 16.766 16.105 15.738 15.894 15.915	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.624 74.595 72.182 72.451 70.963 70.469 69.921 68.988	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 36.64 1.00 36.66 1.00 23.79 1.00 23.94 1.00 38.43 1.00 38.53 1.00 46.42 1.00 39.54 1.00 39.54 1.00 25.87 1.00 42.01 1.00 2.00 1.00 36.68 1.00 36.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3288 3299 32992 32993 32995 32996 32997 33001 33002 3303 3304	CD1 CD2 C O N CAB CG CD OE1 C O N CAB CG CD OE2 C O N CAB CG CD OE2	LEU LEU LEU SER SER SER GLIN GLIN GLIN PRO PRO PRO	547 547 547 548 548 548 549 549 555 555 555 555 555 555 555	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.384 44.7567 43.567 43.560 42.002 46.456 47.634 46.068 44.775 46.253 46.253 46.886 47.816	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.672 20.074 16.547 16.766 16.105 15.738 15.894 15.915 17.184	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 73.624 74.595 72.182 72.451 70.963 70.469 69.921 68.988 69.580	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 46.80 1.00 38.43 1.00 46.42 1.00 39.54 1.00 25.87 1.00 42.01 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 36.68 1.00 36.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3289 3299 3299 3299 3299	CD1 CD2 C O N CA CB CC O N CA CCB CC O N CC CC O N CC C	LEU LEU LEU SER SER SER SER GLIN GLIN PRO	547 547 547 548 548 548 548 549 5549 5555 5555 5555	52.096 52.905 48.422 48.642 47.230 46.091 45.099 45.611 45.374 44.299 45.384 44.7567 43.567 43.568 44.7567 43.934 44.7002 46.456 47.634 46.068 44.729 47.075 46.253 44.886 47.816 47.318	18.701 17.363 16.526 15.414 16.881 15.964 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.074 16.766 16.766 16.705 15.738 15.894 15.350 15.350 15.350 15.350	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.852 75.406 74.569 73.084 73.990 74.054 73.990 74.595 72.182 72.451 70.963 70.469 69.921 68.750 68.988 69.580	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 36.66 1.00 23.79 1.00 23.79 1.00 46.80 1.00 38.43 1.00 38.43 1.00 38.43 1.00 38.53 1.00 46.42 1.00 39.54 1.00 25.87 1.00 42.01 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3289 3290 3299 3299 3299 3299 3299 3299 329	CD1 CD2 CONCACGCCD NCCACGCCD NCCACGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	LEU LEU LEU SER SER SER SER GLIN NG GLIN PRO	547 547 547 548 548 548 549 549 555 555 555 555 555 555 555 555	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.384 44.751 43.567 43.180 43.934 44.729 47.075 46.068 44.729 47.075 46.886 47.816 47.318 49.013	18.701 17.363 16.526 15.414 16.881 15.966 17.389 15.975 15.381 16.649 16.754 18.128 18.354 19.800 20.074 16.766 16.105 15.738 15.894 15.350 15.350 15.350 17.184 18.280 17.046	78.934 77.029 76.434 75.946 76.903 76.993 78.821 75.552 75.406 74.569 73.242 73.084 73.990 74.054 73.624 73	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.19 1.00 36.66 1.00 23.79 1.00 23.79 1.00 46.80 1.00 38.43 1.00 38.43 1.00 38.53 1.00 46.42 1.00 39.54 1.00 25.87 1.00 42.01 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3289 3299 3299 3299 3299	CD1 CD2 CONCABOCONCACONCACONCACONCACONCACONCACON	LEU LEU LEU LEU SER	547 547 547 548 548 548 548 549 5549 5555 5555 5555	52.096 52.905 48.422 48.642 47.230 45.099 45.611 45.374 44.299 45.384 44.751 43.567 43.180 43.934 42.002 46.453 44.729 47.075 46.253 44.886 47.318 47.318 47.318	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 18.354 19.800 20.672 20.074 16.547 16.738 15.915 17.184 15.915 17.184 18.280 17.046 18.198	78.934 77.029 76.434 75.946 76.903 76.993 78.821 75.552 75.406 74.569 73.242 73.990 74.054 73.624 74.595 72.451 70.469 69.921 68.750 68.988 69.580 69.015 68.626	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 23.94 1.00 38.53 1.00 46.80 1.00 38.53 1.00 46.42 1.00 39.54 1.00 39.54 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3289 3299 3299 3299 3299	CD1 CD2 CONCABCCONCABCCONCABCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCC	LEU LEU LEU LEU SER	547 547 547 548 548 5548 5548 5549 5555555555555555	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.968 45.384 44.751 43.567 43.180 43.934 42.002 46.47.634 46.068 47.634 46.053 44.729 47.075 46.253 44.886 47.318 47.318 49.824 51.293	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 19.800 20.672 20.074 16.547 16.766 16.766 15.350 15.915 17.184 18.280 17.046 18.198 17.788	78.934 77.029 76.434 75.946 76.903 76.898 77.893 78.821 75.552 75.406 74.569 73.242 73.990 74.054 73.624 74.595 72.182 72.451 70.469 69.921 68.750 68.988 69.840 69.840 69.840 69.840 69.840 69.840 69.840	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.64 1.00 36.66 1.00 23.79 1.00 23.94 1.00 38.43 1.00 38.43 1.00 38.53 1.00 46.42 1.00 39.54 1.00 25.87 1.00 42.01 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3289 3299 3299 3299 3299	CD12 CONCABGCONACCONCABGCONACCGCCONACCGCCCCCCCCCCCCCCCCCCCCCCCC	LEU LEU LEU LEU SER	547 547 547 548 548 548 5548 5549 5555 5555 5555 55	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.968 45.384 44.3567 43.180 43.934 42.002 46.456 47.634 46.456 47.675 46.253 44.886 47.818 49.824 51.293 51.490	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 19.800 20.672 20.074 16.547 16.765 15.738 15.894 15.350 15.915 17.184 18.280 17.048 17.788 16.914	78.934 77.029 76.434 75.946 76.903 76.898 77.993 78.821 75.552 75.406 74.569 73.242 73.084 74.054 74.595 72.182 72.451 70.963	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 36.19 1.00 39.64 1.00 36.66 1.00 23.79 1.00 23.79 1.00 23.94 1.00 38.53 1.00 46.80 1.00 38.53 1.00 46.42 1.00 39.54 1.00 39.54 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3279 3280 3281 3282 3288 3288 3289 3299 3299 3299 3299	CD1 CD2 CONCABCCONCABCCONCABCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCCONCABCCC	LEU LEU LEU LEU SER	547 547 547 548 548 5548 5548 5549 5555555555555555	52.096 52.905 48.422 48.642 47.230 46.099 45.611 45.374 44.299 45.968 45.384 44.751 43.567 43.180 43.934 42.002 46.47.634 46.068 47.634 46.053 44.729 47.075 46.253 44.886 47.318 47.318 49.824 51.293	18.701 17.363 16.526 15.414 16.881 15.964 16.356 17.389 15.975 15.381 16.649 16.754 19.800 20.672 20.074 16.547 16.766 16.766 15.350 15.915 17.184 18.280 17.046 18.198 17.788	78.934 77.029 76.434 75.946 76.903 76.898 77.893 78.821 75.552 75.406 74.569 73.242 73.990 74.054 73.624 74.595 72.182 72.451 70.469 69.921 68.750 68.988 69.840 69.840 69.840 69.840 69.840 69.840 69.840	1.00 16.17 1.00 11.72 1.00 2.00 1.00 6.91 1.00 5.34 1.00 36.64 1.00 36.66 1.00 23.79 1.00 23.94 1.00 38.43 1.00 38.53 1.00 46.80 1.00 39.54 1.00 39.54 1.00 25.87 1.00 42.01 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 36.68 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000

MOTA	3315	CD1 I		51	53.648	18.701	68.46B	1.00	2.00	0
MOTA	3316	_		51	49.295 49.596	18.705 19.807	67.281 66.840	1.00	2.00	ŏ
MOTA	3317	-		551 552	48.514	17.863	66.627	1.00	2.00	Ö
MOTA	3318 3320			552	47.896	18.202	65.369	1.00	2.00	0
MOTA MOTA	3321			552	48.287	17.148	64.320	1.00	2.00	0
ATOM	3322			552	47.809	17.215	62.867	1.00	2.00	0
ATOM	3323	CD1 L		552	48.082	18.532	62.209	1.00	2.00	0
MOTA	3324		_	552	48.544	16.174 18.180	62.131 65.728	1.00	2.00	0
MOTA	3325			552 552	46.397 45.753	17.124	65.734	1.00	2.00	ŏ
MOTA	3326			553	45.881	19.352	66.100	1.00	20.15	0
MOTA MOTA	3327 3329			553	44.485	19.512	66.517		20.15	0
ATOM	3330		BU S	553	44.240	20.926	67.055	1.00	2.00	0
ATOM	3331			553	44.374	21.262	68.543 69.290	1.00	2.00 2.00	0
MOTA	3332	CD1 L	_	553	45.224	20.282 22.639	68.638	1.00	2.00	ő
MOTA	3333			553 5 53	44.942 43.495	19.229	65.410		20.15	ŏ
MOTA	3334 3335	-		553	43.604	19.780	64.309	1.00	2.00	Õ
MOTA MOTA	3336	_		554	42.524	18.375	65.724		14.62	0
ATOM	3338			554	41.476	17.983	64.786		13.80	0
ATOM	3339		LU :	554	41.135	16.492	64.956		57.31	0
MOTA	3340	-		554	42.326	15.550	65.177 63.927		69.83 73.58	0
MOTA	3341			554 FF4	43.168 42.918	15.310 15.957	62.887		82.45	ŏ
ATOM	3342	OE1 G		554 554	44.090	14.465	63.988		79.54	0
MOTA MOTA	3343 3344	_		554	40.241	18.823	65.105		12.33	0
ATOM	3345	-		554	39.253	18.311	65.638		51.99	0
MOTA	3346	N I		555	40.296	20.112	64.792	1.00	2.00	0
MOTA	3348			555	39.176	21.007 22.456	65.077 64.911	1.00	2.00	ő
MOTA	3349			555 	39.628 40.660	22.840	65.972	1.00	2.00	ŏ
ATOM	3350	CG I		555 555	41.274	24.194	65.697	1.00	2.00	0
MOTA MOTA	3351 3352	CD2 I		555	39.959	22.839	67.294	1.00	2.00	0
ATOM	3353			555	37.932	20.734	64.237	1.00	2.00	0
MOTA	3354	0 1		555	37.862	19.731	63.537	1.00	2.00	0
MOTA	3355			556	36.933	21.603	64.345 63.569	1.00	2.00	ő
MOTA	3357			556 556	35.716 35.028	21.460 20.115	63.846	1.00		ō
MOTA	3358	-		556 556	34.175	20.063	65.100		48.42	0
MOTA MOTA	3359 3360			556	32.935	19.176	64.941		58.49	0
ATOM	3361			556	32.792	18.188	65.711		60.59	0
ATOM	3362	OE2 G		556	32.101	19.477	64.047	1.00	59.64 2.00	0
MOTA	336 3			556	34.736	22.583 23.580	63.826 64.44 6	1.00	16.83	Ö
MOTA	3364			556 557	35.067 33.544	22.423	63.257	1.00	2.00	0
MOTA	3365 3367			557	32.398	23.330	63.427	1.00	2.00	0
MOTA MOTA	3368	-		557	31.747	23.013	64.788	1.00		0
ATOM	3369		ALA	557	32.701	24.826	63.286	1.00	2.00	0
MOTA	3370	0 1		557	33.711	25.191 25.711	62.671 63.802		40.85	Ö
MOTA	3371			558	31.811 30.459	25.711 25.558	64.353	1.00	2.44	Ö
MOTA	3372			558 558	32.123	27.139	63.658	1.00	2.00	0
MOTA MOTA	3373 3374			558	30.786	27.826	63.951	1.00	2.53	0
MOTA	3375			558	29.798	26.759	63.843	1.00		0
MOTA	3376		-	558	33.190	27.648	64.627	1.00	2.00 7.53	Ö
MOTA	3377			558	32.975	27.616 28.119	65.844 64.112	1.00	2.00	ő
MOTA	3378			559	34.325 35.349	28.689	64.982	1.00	2.00	Ō
MOTA	3380			559 5 5 9	36.568	27.769	65.148	1.00	2.00	0
MOTA MOTA	3381 3382			559	37.592	27.571	64.050	1.00	2.00	0
MOTA	3383	CD1		559	38.620	26.563	64.476	1.00	2.00	0
MOTA	3384	CD2		559	36.885	27.083	62.830	1.00	2.00	0
MOTA	3385		LEU	559	35.769	30.038 30. 38 9	64.435 63.311	1.00	2.00	0
MOTA	3386		LEU	559 560	35.444 36.461	30.389 30.804	65.260	1.00	2.00	ő
MOTA	3387		LYS	560 560	36.461	32.115	64.889	1.00	2.00	Ö
ATOM	33 8 9 3390		LYS LYS	560	36.359	33.143	65.843		20.44	0
ATOM	חברנ	C12 1	~ · · ·							

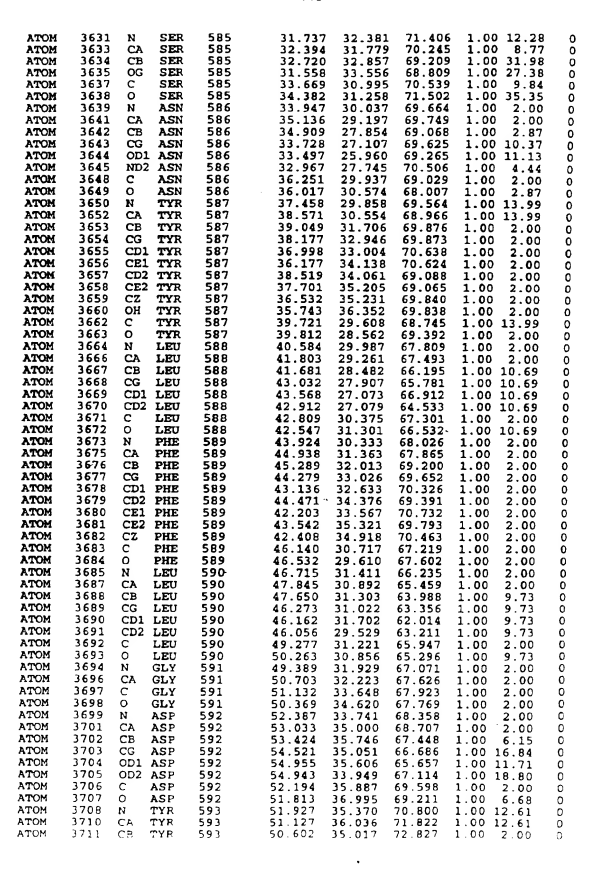
ATOM	3391	CG	LYS	560	36.612	34.562	65.427	1.00 20.44	_
									0
MOTA	3392	CD	LYS	560	35.574	35.512	66.018	1.00 20.44	0
ATOM	3393	CE	LYS	560	34.253	35.452	65.278	1.00 20,44	0
ATOM	3394	NZ	LYS	560	33.706	34.083	65.225		
								1.00 20.44	0
ATOM	3398	С	LYS	560	38.426	31.990	65.052	1.00 2.00	0
MOTA	3399	0	LYS	560	38.908	31.624	66.117	1.00 20.44	
									0
ATOM	3400	N	ILE	561	39.164	32.244	63.986	1.00 2.00	0
ATOM	3402	CA	ILE	561	40.606	32.106	64.017	1.00 2.00	ō
				561					
MOTA	3403	CB	ILE		41.095	31.318	62.750	1.00 2.00	0
MOTA	3404	CG2	ILE	561	42.566	30.913	62.891	1. 0 0 2.00	0
MOTA	3405	CG1	ILE	561	40.204	30.076	62.558	1.00 2.00	ŏ
MOTA	3406	CD1	ILE	561	40.806	28.940	61.761	1.00 2.00	0
MOTA	3407	С	ILE	561	41.247	33.488	64.102	1.00 2.00	0
ATOM	3408	ō	ILE	561	40.703	34.450	63.543		
									0
MOTA	3409	N	CYS	562	42.365	33.586	64.833	1.00 2.00	0
ATOM	3411	CA	CYS	562	43.124	34.829	65.001	1.00 2.00	0
				562	42.793				
MOTA	3412	CB	CYS			35.464	66.339	1.00 13.31	0
ATOM	3413	SG	CYS	562	41.073	35.733	66.574	1.00 13.31	0
MOTA	3414	C	CYS	562	44.642	34.589	64.928		
									0
MOTA	3415	0	CYS	562	45.151	33.551	65.373	1.00 13.31	0
ATOM	3416	N	GLY	563	45.362	35.540	64.348	1.00 2.00	0
MOTA	3418	CA	GLY	563	46.805	35.424	64.252	1.00 2.00	0
ATOM	3419	С	GLY	563	47.546	36.260	65.291	1.00 2.00	0
MOTA	3420	ŏ	GLY	563	46.997	36.556	66.351		
									0
MOTA	3421	N	ASP	564	48.780	36.649	64.958	1.00 2.00	0
ATOM	3423	CA	ASP	564	49.657	37.452	65.811	1.00 2.00	0
MOTA	3424	CB	ASP	564	50.705	38.191	64.964	1.00 2.00	0
MOTA	3425	CG	ASP	564	51.724	37.270	64.321	1.00 2.00	0
ATOM	3426		ASP	564	51.390	36.585	63.334	1.00 2.00	ŏ
MOTA	3427	OD2	ASP	564	52.876	37.249	64.786	1.00 2.00	0
ATOM	3428	С	ASP	564	48.981	38.497	66.693	1.00 2.00	0
				564	48.395	39.442	66.189		
MOTA	3429	0	ASP						0
ATOM	3430	N	ILE	56 5	49.094	38.327	68.009	1.00 31.53	0
MOTA	3432	CA	ILE	565	48.537	39.294	68.953	1.00 35.54	0
	-		-						
MOTA	3433	CB	ILE	56 5	47.810	38.612	70.135	1.00 9.39	0
ATOM	3434	CG2	ILE	56 5	47.139	39.664	70.9 9 6	1.00 9.39	0
MOTA	3435		ILE	565	46.710	37.685	69.622	1.00 9.39	0
		CG1							
ATOM	3436	CD1	ILE	56 5	45.611	38. 424	68.906	1.00 9.39	0
MOTA	3437	С	ILE	565	49.681	40.168	69.489	1.00 32.16	0
								1.00 9.39	ō
MOTA	3438	0	ILE	565	49.491	41.356	69.756		
ATOM	3439	N	HIS	56 6	50.859	39.560	69.637	1.00 14.33	0
MOTA	3441	CA	HIS	566	52.082	40.219	70.111	1.00 15.67	0
		_				41.227	71.245		ŏ
MOTA	3442	С	HIS	566	51.958			1.00 16.54	-
MOTA	3443	0	HIS	566	52.427	42.359	71.127	1.00 15.89	0
MOTA	3444	CB	HIS	56 6	52.811	40.882	68.943	1.00 9.53	0
								1.00 9.53	
MOTA	3445	CG	HIS	566	53.652	39.936	6B.153		0
ATOM	3446	ND1	HIS	56 6	54.829	39.391	68.605	1.00 9.53	0
MOTA	3448	CD2	HIS	566	53.470	39.432	66,910	1.00 9.53	0
				_					ō
MOTA	3449	NEZ	HIS	566	5 4.52 5	38.582	66.589		
MOTA	3450	CE1	HIS	566	55. 31 2	38. 60 0	67.652	1.00 9.53	0
ATOM	3451	N	GLY	567	51.337	40.808	72.345	1.00 2.00	0
MOTA	3453	CA	GLY	567	51.170	41.686	73.490	1.00 2.00	0
MOTA	3454	С	GLY	567	50.249	42.894	73.375	1.00 2.00	0
MOTA	3455			567	50.269	43.764	74.254	1.00 3.59	0
		0	GLY						
ATOM	3456	N	GLN	568	49.455	42.973	72.314	1.00 36.66	0
MOTA	3458	CA	GLN	568	48.527	44.086	72.141	1.00 34.60	0
							70.667	1.00 2.50	0
MOTA	3459	CB	GLN	568	48.164	44.236			
MOTA	3460	CG	GLN	568	49.345	44.164	69.715	1.00 3.37	0
MOTA	3461	CD	GLN	568	49.768	45.507	69.144	1.00 .3.10	0
ATOM	3462	OEI	GLN	568	50.958	45.769	68.974	1.00 4.87	0
MOTA	3463	NE2	GLN	568	48.800	46.349	68.813	1.00 18.91	0
ATOM	3463				47.280	43.730	72.950	1.00 34.26	0
AIUN	3463		CT N					A . U U J W . A U	
	3466	C	GLN	568					
MOTA			GLN GLN	568	46.253	43.365	72.380	1.00 7.33	0
	3466 3467	C 0	GLN	568		43.365			
MOTA	3466 3467 3468	0 0	GLN TYR	568 569	46.253 47.363	43.365 43.864	72.380 74.272	1.00 7.33 1.00 2.00	0
MOTA MOTA	3466 3467 3468 3470	C O N CA	GLN TYR TYR	568 569 569	46.253 47.363 46.263	43.365 43.864 43.485	72.380 74.272 75.161	1.00 7.33 1.00 2.00 1.00 2.00	0
MOTA	3466 3467 3468	0 0	GLN TYR	568 569 569 569	46.253 47.363 46.263 46.664	43.365 43.864 43.485 43.687	72.380 74.272 75.161 76.626	1.00 7.33 1.00 2.00 1.00 2.00 1.00 26.93	0 0 0
MOTA MOTA MOTA	3466 3467 3468 3470 3471	C O N CA CB	GLN TYR TYR TYR	568 569 569 569	46.253 47.363 46.263	43.365 43.864 43.485	72.380 74.272 75.161	1.00 7.33 1.00 2.00 1.00 2.00	0
ATOM ATOM ATOM AOTA	3466 3467 3468 3470 3471 3472	C O N CA CB CG	GLN TYR TYR TYR TYR	568 569 569 569	46.253 47.363 46.263 46.664 45.692	43.365 43.864 43.485 43.687 43.048	72.380 74.272 75.161 76.626 77.600	1.00 7.33 1.00 2.00 1.00 2.00 1.00 26.93 1.00 27.02	0 0 0 0
MOTA MOTA MOTA	3466 3467 3468 3470 3471	C O N CA CB	GLN TYR TYR TYR TYR	568 569 569 569	46.253 47.363 46.263 46.664	43.365 43.864 43.485 43.687	72.380 74.272 75.161 76.626	1.00 7.33 1.00 2.00 1.00 2.00 1.00 26.93	0 0 0

							70 303		24 12	_
MOTA	3474	CE1 T	MYR	569	44.496	41.097	78.393	1.00	24.12	0
				569	45.049	43.807	78.575	1.00	24.68	0
ATOM	3475		ľYR							
ATOM	3476	CE2 T	ryr	569	44.139	43.223	79.449		24.01	0
			TYR .	569	43.866	41.869	79.352	1.00	28.35	0
ATOM	3477	_		-						
MOTA	3478	он т	ryr	569	42.950	41.282	80.197	1.00	35.43	0
	-				44.888	44.085	74.946	1.00	2.00	0
MOTA	3480	CI	ryr	569						
ATOM	3481	0 7	MYR	569	43.885	43.374	75.015	1.00	29.07	0
ATOM		-					74.710	1.00	11.04	0
MOTA	3482	N I	ryr	57 0	44.817	45.384				
	3484	CA T	ryr	570	43.515	45.999	74.510	1.00	14.25	0
ATOM		_					74.766	1 00	45.12	0
MOTA	3485	CB 1	ľYR	570	43.599	47.505				
		CG 7	ľYR	570	43.577	47.792	76.261	1.00	51.43	0
ATOM	3486						77.025		48.11	
ATOM	3487	CD1 T	ryr -	570	44.749	47.743				0
			TYR .	570	44.715	47.952	78.398	1.00	52.04	0
MOTA	3488									
ATOM	3489	CD2 1	TYR .	570	42.375	48.063	76.917		49.50	0
				570	42.340	48.272	78.277	1 00	51.48	0
MOTA	3490		TYR							
ATOM	3491	CZ 1	ľYR	570	43.507	48.215	79.010	1.00	53.24	0
					43.452	48.426	80.359	1 00	54.97	0
MOTA	3492	OH 7	ryr	57 0						
ATOM	3494	C 7	ry r	570	42.929	45.653	73.152	1.00	13.58	0
					41.708	45.654	72.967	1 00	43.14	0
MOTA	3495	0 7	ryr	570	41.700					
	3496	N A	ASP	571	43.813	45.311	72.217	1.00	12.28	0
MOTA										
MOTA	3498	CA A	ASP	571	43.402	44.903	70.891	1.00	11.58	0
			ASP	571	44.590	44.870	69.975	1.00	6.85	0
MOTA	3499									
MOTA	3500	CG 2	ASP	571	45.128	46.237	69.733	1.00	12.88	0
				T	46.254	46.522	70.185	1.00	19.21	0
MOTA	3501	OD1 A	ASP	571						
MOTA	3502	OD2 A	ACP	571	44.402	47.044	69.110	1.00	16.01	0
						43.539	71.062	1 00	14.93	0
MOTA	3503	C 2	ASP	571	42.818	43.333				
	3504	0 2	ASP	571	41.775	43.247	70.507	1.00	19.59	0
MOTA		_				40 707		1.00	2.00	0
MOTA	3505	N I	LEU	572	43.487	42.707	71.853			
			LEU	572	42.977	41.378	72.147	1.00	2.00	0
MOTA	3507								2.00	
MOTA	3508	CB I	LEU	572	43.909	40.641	73.110	1.00	2.00	0
				572	43.302	39.416	73.820	1.00	2.00	0
ATOM	3509	CG I	LEU							
ATOM	3510	CD1 I	JRU	572 -	43.061	38.267	72.832	1.00	2. 0 0	0
		-			44.227	38.982	74.955	1.00	2.00	0
ATOM	3511	CD2 I	LEU	572						
MOTA	3512	C I	LEU	572	41.579	41.554	72.771	1.00	2.00	0
							72.512	1.00	2.00	0
ATOM	3513	0 1	LEU	572	40.678	40.758				
		_	LEU	573	41.395	42.604	73.572	1.00	9.95	0
atom	3514								6.25	0
MOTA	3516	CA I	LEU	573	40.092	42.874	74.195	1.00		
				573	40.224	43.910	75.317	1.00	2.00	0
MOTA	3517		LEU							0
MOTA	3518	CG I	LEU	573	40.807	43.360	76.625	1.00	2.00	
		CD1 I		573	40.667	44.416	77.717	1.00	2.00	0
MOTA	3519							1.00	2.00	
MOTA	3520	CD2 I	LEU	573	40.079	42.053				0
							77.033	1.00	2.00	0
MOTA	3521				39 NSS					U
ATOM		C I	LEU	573	39.055	43.344	73.170	1.00	4.08	
	3522	-								0
	3522	0 1	LEU	573	37.865	43.344 42.974	73.170 73.260	1.00 1.00	4.08	0
ATOM	3522 3523	0 1			37.865 39.518	43.344 42.974 44.153	73.170 73.260 72.203	1.00 1.00 1.00	4.08 2.00 13.12	0
ATOM	3523	0 I	LEU ARG	573 574	37.865 39.518	43.344 42.974 44.153	73.170 73.260	1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30	0
MOTA MOTA	3523 3 52 5	O I	LEU ARG ARG	573 574 574	37.865 39.518 38.682	43.344 42.974 44.153 44.665	73.170 73.260 72.203 71.115	1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30	0 0 0
ATOM	3523	O I	LEU ARG	573 574	37.865 39.518	43.344 42.974 44.153 44.665 45.598	73.170 73.260 72.203 71.115 70.212	1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04	0000
MOTA MOTA	3523 3525 3526	O I N J CA J CB J	LEU ARG ARG ARG	573 574 574 574	37.865 39.518 38.682 39.491	43.344 42.974 44.153 44.665 45.598	73.170 73.260 72.203 71.115 70.212	1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30	0 0 0
ATOM ATOM ATOM MOTA	3523 3525 3526 3527	O I N J CA J CB J CG J	LEU ARG ARG ARG ARG	573 574 574 574 574	37.865 39.518 38.682 39.491 39.704	43.344 42.974 44.153 44.665 45.598 46.994	73.170 73.260 72.203 71.115 70.212 70.757	1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00	0 0 0 0
ATOM ATOM ATOM MOTA	3523 3525 3526 3527	O I N J CA J CB J CG J	LEU ARG ARG ARG	573 574 574 574	37.865 39.518 38.682 39.491	43.344 42.974 44.153 44.665 45.598	73.170 73.260 72.203 71.115 70.212 70.757 69.915	1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07	0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528	O II N II CA II CB II CG II CD II	LEU ARG ARG ARG ARG ARG	573 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697	43.344 42.974 44.153 44.665 45.598 46.994 47.811	73.170 73.260 72.203 71.115 70.212 70.757 69.915	1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07	0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529	O II N II CA II CB II CG II CD II NE II	LEU ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.989	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514	1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10	000000
ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528	O II N II CA II CB II CG II CD II NE II	LEU ARG ARG ARG ARG ARG	573 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697	43.344 42.974 44.153 44.665 45.598 46.994 47.811	73.170 73.260 72.203 71.115 70.212 70.757 69.915	1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14	0000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531	O II N II CA II CB II CG II CD II NE II CZ II	LEU ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.989 48.625	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587	1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14	000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532	O I N I I CA I I CB I	LEU ARG ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.989 48.625 49.162	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84	00000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532	O II N II CA II CB II CG II CD II NE II CZ II	LEU ARG ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.989 48.625	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84 40.46	000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535	O II N II CA II CB II CG II CD II NE II CZ II NH1 II NH2 II	LEU ARG ARG ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.881 47.881 47.811 48.625 49.162 48.713	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84 40.46	000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538	O I I I I I I I I I I I I I I I I I I I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 38.265	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.989 48.625 49.162 48.713 43.440	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.890 66.341 70.324	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99	00000000000
ACOM ACOM ACOM ACOM ACOM ACOM ACOM ACOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538	O I I I I I I I I I I I I I I I I I I I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG	573 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.881 47.881 47.811 48.625 49.162 48.713	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84 40.46	000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3539	O II N I I CA I I CB I C	LEU ARG	573 574 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 40.565 38.265 37.092	43.344 42.974 44.153 44.665 45.598 46.994 47.811 47.989 48.625 49.162 48.713 43.440 43.253	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.587 66.341 70.324 70.004	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16	00000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3539 3540	O I I N I I CA I I CB	LEU ARG	573 574 574 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 38.265 37.092 39.260	43.344 42.974 44.153 44.665 45.598 46.5994 47.811 47.989 48.625 49.162 48.713 43.440 43.253 42.591	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.066	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 40.46 10.99 30.16 2.00	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3539 3540	O I I N I I CA I I CB I	LEU ARG	573 574 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 40.565 38.265 37.092	43.344 42.974 44.153 44.665 45.598 467.811 47.989 48.625 49.162 48.713 43.445 43.2591 41.339	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 40.46 10.99 30.16 2.00 2.00	0000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3539 3540 3542	O I I CA	LEU ARG	573 574 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 37.092 39.260 39.156	43.344 42.974 44.153 44.665 45.598 467.811 47.989 48.625 49.162 48.713 43.445 43.2591 41.339	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 40.46 10.99 30.16 2.00 2.00	0000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3540 3542 3543	O II N I CB II CB	LEU ARG	573 574 574 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 38.265 37.092 39.156 40.471	43.344 42.974 44.153 44.665 45.598 46.981 47.989 48.625 49.162 48.713 43.253 42.591 40.578	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.0066 69.323 69.502	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84 40.46 10.99 30.16 2.00 2.00 18.14	00000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3540 3542 3543	O II N I CB II CB	LEU ARG	573 574 574 574 574 574 574 574 574 574 574	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 38.265 37.092 39.156 40.471	43.344 42.974 44.153 44.665 45.598 46.981 47.989 48.625 49.162 48.713 43.253 42.591 40.578	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 40.46 10.99 30.16 2.00 2.00	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3532 3535 3535 3540 3544 3544	O I I N I CB I I CB I I CB I CC I CC I CC	LEU ARG	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 38.265 37.092 39.260 39.156 40.471 41.058	43.344 42.974 44.153 44.665 45.598 46.9811 47.989 48.625 49.162 48.713 43.445 43.2591 41.339 40.578 39.569	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323 69.502 68.514	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 3531 3532 3535 3538 3540 3542 3543	O II N I CB II CB	LEU ARG	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 37.092 39.265 39.265 39.265 40.471 41.058 40.112	43.344 42.974 44.153 44.665 45.598 46.981 47.819 48.625 49.162 48.713 43.253 42.531 41.339 42.531 41.3378 39.569 38.378	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.066 69.323 69.502 68.514 68.370	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14	0000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3532 3535 3535 3540 35442 35443 35444 3545	O I I I CA I I CA I CCA	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 37.092 39.265 39.265 39.265 40.471 41.058 40.112	43.344 42.974 44.153 44.665 45.598 46.981 47.819 48.625 49.162 48.713 43.253 42.531 41.339 42.531 41.3378 39.569 38.378	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323 69.502 68.514	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3532 3535 3535 3544 3544 3544 3544 3546	O I I I C I I C C I I I C C I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 37.092 39.265 39.265 39.265 40.471 41.058 40.112 41.355	43.344 42.974 44.153 44.665 45.598 46.981 47.819 48.625 49.162 48.713 43.440 43.253 42.531 41.3378 39.569 38.378 40.249	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.006 69.323 69.323 69.323 68.514 68.370 67.176	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3532 3535 3535 3540 35442 35443 35444 3545	O I I I C I I C C I I I C C I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 37.092 39.265 39.265 39.265 40.471 41.058 40.112	43.344 42.974 44.153 44.665 45.598 46.981 47.819 48.625 49.162 48.713 43.253 42.531 41.339 42.531 41.3378 39.569 38.378	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3535 3535 3535 3540 35443 35444 35446 3547	O I I I I I I I I I I I I I I I I I I I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 37.092 39.156 40.471 41.058 40.112 41.355 37.970	43.344 42.974 44.153 44.665 45.598 46.991 47.819 48.625 49.162 48.713 43.253 43	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3535 3535 3535 3540 35443 35444 35446 3547 3548	O I I I CA I I C	LEU ARG	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 37.092 39.260 39.156 40.471 41.058 40.112 41.355 37.970 37.121	43.344 42.974 44.153 44.665 45.598 46.991 47.819 47.9625 49.1623 43.4453 43.42.591 41.5769 40.5769 40.494 40.081	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.890 66.341 70.324 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801 69.001	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3531 3535 3535 3535 3540 35443 35444 35446 3547 3548	O I I I CA I I C	LEU ARG	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 42.198 40.565 37.092 39.156 40.471 41.058 40.112 41.355 37.970	43.344 42.974 44.153 44.665 45.598 46.991 47.819 48.625 49.162 48.713 43.253 43	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.324 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14 2.00 18.14 29.25	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 3525 3526 3527 3528 3529 35332 35335 35340 35443 35444 35445 35445 35446 35448 3549	O I I I C I I C C I I I C C I I I C C I I I C C I I I C C I I I C C I I I C C I I I C C I I I C C I I I I C C I I I I C C I I I I C C I I I I C C I I I I C C I I I I C C I I I I C C I I I I C I I I I C I I I I C I I I I I C I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU LEU PHE	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 40.565 37.092 39.156 40.471 41.058 40.112 41.355 37.970 37.121 37.908	43.344 42.974 44.153 44.6658 45.5984 47.9891 47.989 48.6252 48.713 43.2491 40.578 39.569 38.378 40.249 40.081 40.247	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801 69.801 71.103	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14 2.00 18.14 29.25	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 355229 355335 355335 355340 355443 355446 355446 355446 355446 35546 35541	O I I I C I I C C I I I C C I I I I C C I I I I C I I I C I I I C I I I C I I I C I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I I C I	LEU ARG	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 37.092 39.156 40.471 41.058 40.112 41.355 37.970 37.121 37.908 36.823	43.344 42.974 44.153 44.6658 45.5984 47.989 48.625 49.162 48.713 43.42591 41.339 40.578 39.578 40.249 40.247 39.459	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801 69.801 71.103 71.664	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 36.14 35.84 40.46 10.99 30.16 2.00 18.14 18.14 18.14 18.14 18.14 29.25 32.60	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 355229 355335 355335 355340 355443 355446 355446 355446 355446 35546 35541	O I I I C I I C C I I I C C I I I I C C I I I I C I I I C I I I C I I I C I I I C I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I I C I	LEU ARG	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.697 40.294 41.018 40.565 37.092 39.156 40.471 41.058 40.112 41.355 37.970 37.121 37.908	43.344 42.974 44.153 44.6658 45.5984 47.9891 47.989 48.6252 48.713 43.2491 40.578 39.569 38.378 40.249 40.081 40.247	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801 69.801 71.103	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14 2.00 18.14 29.25	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 35229 35335 35335 35335 35344 35446 35447 355447 355447 35552	O I I I C I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I I C I I I I I C I I I I I C I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU PHE PHE	573 574 574 574 574 574 574 574 574 575 575	37.865 39.518 38.682 39.491 39.704 40.294 41.018 42.198 40.565 37.092 39.156 40.112 41.355 37.970 37.121 37.908 36.823 37.115	43.344 42.974 44.153 44.665 45.598 467.989 47.989 48.625 49.162 48.1259 40.259 40.578 39.569 40.249 40.0817 39.4098	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.0066 69.323 69.502 68.514 68.370 67.176 69.801 71.1664 73.119	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14 18.14 2.00 18.14 29.25 32.60 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 355229 355335 355335 355340 355443 355446 355446 355446 355446 35546 35541	O I I I C I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I I C I I I I I C I I I I I C I	LEU ARG	573 574 574 574 574 574 574 574 575 575 575	37.865 39.518 38.682 39.491 40.294 40.294 41.018 42.198 40.565 37.092 39.156 40.112 41.355 37.121 37.908 37.121 37.908 37.115 38.115	43.344 42.974 44.153 44.665 45.598 467.989 47.989 48.625 49.162 48.713 43.259 40.259 40.578 39.569 840.249 40.081 40.081 40.081 40.081 39.098 39.098	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801 71.103 71.664 73.119 73.270	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.07 36.10 35.84 40.46 10.99 30.16 2.00 18.14 18.14 18.14 18.14 18.14 2.00 18.14 2.00 2.00 2.00 2.00 2.00 2.00 2.00 2.0	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 35229 355335 355335 3553442 355447 355447 355447 355447 35553	O I I I C I I I C I I I C I I I C I I C I I I C I I I C I I I C I I I C I I I I C I I I C I I I C I I I C I I I C I I I C I I I C I I I C I I I C I I I C I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I C I I I I I C I I I I I C I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU LEU PHE PHE	573 574 574 574 574 574 574 574 575 575 575	37.865 39.518 38.682 39.491 40.294 40.294 41.018 42.198 40.565 37.092 39.156 40.112 41.355 37.121 37.908 37.121 37.908 37.115 38.115	43.344 42.974 44.153 44.665 45.598 467.989 47.989 48.625 49.162 48.1259 40.259 40.578 39.569 40.249 40.0817 39.4098	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.006 69.323 68.514 68.370 67.176 69.801 71.1664 73.119 73.270 74.184	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.00 35.07 36.10 35.84 40.46 10.99 30.16 2.00 2.00 18.14 18.14 18.14 18.14 18.14 2.00 18.14 29.25 32.60 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 35228 355335 35535 35535 35534 35544 35544 35555 35555 35554	O I I I CA I CCA I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU PHE PHE PHE	573 574 574 574 574 574 574 574 575 575 575	37.865 39.518 38.682 39.491 39.709 40.294 40.294 41.298 40.565 37.265 39.1566 40.4058 40.4058 40.112 41.355 37.970 37.121 37.908 36.825 37.920 39.155	43.344 42.974 44.1665 45.598 46.8919 47.8985 49.713 43.440.3531 40.2531 40.2531 40.2458 40.	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.006 69.323 68.514 68.370 67.176 69.801 71.1664 73.119 73.270 74.184	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.07 36.10 35.84 40.46 10.99 30.16 2.00 18.14 18.14 18.14 18.14 18.14 2.00 18.14 2.00 2.00 2.00 2.00 2.00 2.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3523 35226 35227 35229 355335 355335 3553442 355447 355447 355447 355447 35553	O I I I I I I I I I I I I I I I I I I I	LEU ARG ARG ARG ARG ARG ARG ARG ARG ARG LEU LEU LEU LEU LEU LEU PHE PHE	573 574 574 574 574 574 574 574 575 575 575	37.865 39.518 38.682 39.491 40.294 40.294 41.018 42.198 40.565 37.092 39.156 40.112 41.355 37.121 37.908 37.121 37.908 37.115 38.115	43.344 42.974 44.153 44.665 45.598 467.989 47.989 48.625 49.162 48.713 43.259 40.259 40.578 39.569 840.249 40.081 40.081 40.081 40.081 39.098 39.098	73.170 73.260 72.203 71.115 70.212 70.757 69.915 68.514 67.587 67.890 66.341 70.004 70.066 69.323 69.502 68.514 68.370 67.176 69.801 71.103 71.664 73.119 73.270	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	4.08 2.00 13.12 12.30 30.04 31.07 36.10 35.84 40.46 10.99 30.16 2.00 18.14 18.14 18.14 18.14 18.14 2.00 18.14 2.00 2.00 2.00 2.00 2.00 2.00 2.00 2.0	000000000000000000000000000000000000000

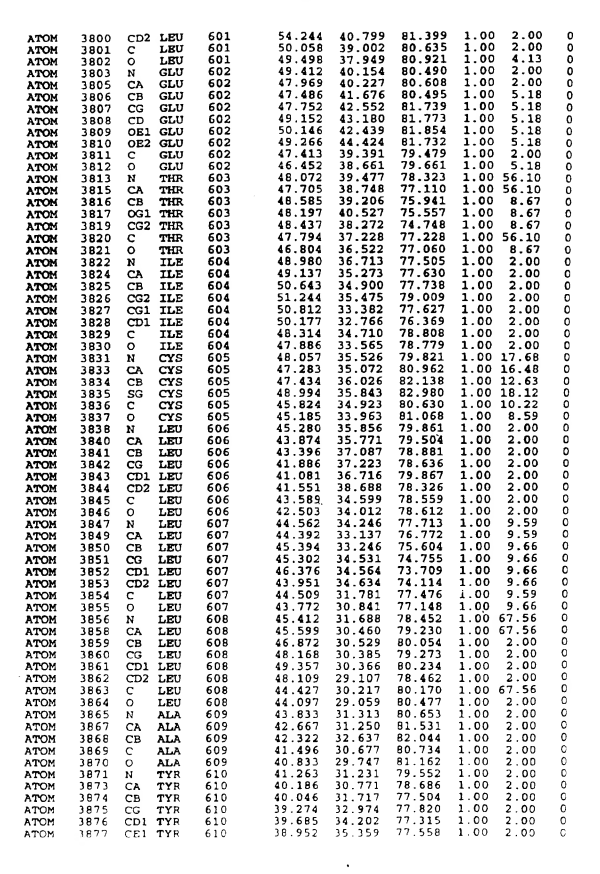
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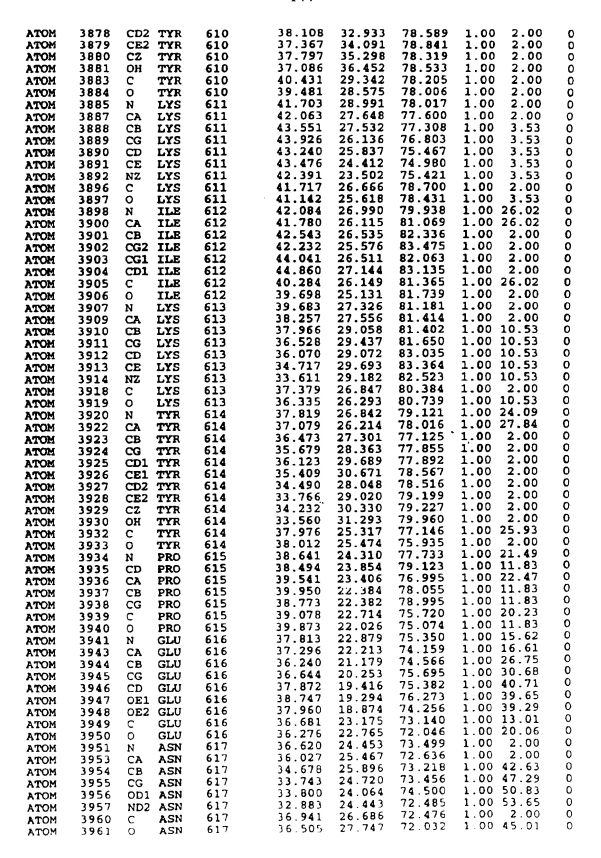


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ATOM 3713 CD1 TYR 593 59.58 32.532 72.382 1.00 2.00 0 0 0 0 0 0 0 0 0	MOTA	3712	CG	TYR	59 3		49.994	33.790	72.200	1.00 2.00	0
ATOM 3714 CEI TTR 593 49,995 31,401 71,793 1.00 2.00 0 ATOM 3716 CEZ TTR 593 48,852 33,886 71,413 1.00 2.00 0 ATOM 3717 CZ TTR 593 48,852 31,860 71,003 2.00 0 ATOM 3718 OH TTR 593 48,855 31,540 31,540 72,560 1.00 2.00 0 ATOM 3720 C TTR 593 51,995 37,019 72,560 1.00 2.00 0 ATOM 3721 O TTR 593 51,995 37,019 72,560 1.00 2.00 0 ATOM 3721 O TTR 593 51,995 37,019 72,560 1.00 2.00 0 ATOM 3722 N VAL 594 51,270 36,660 72,692 1.00 2.00 0 ATOM 3725 CB VAL 594 54,238 37,643 73,259 1.00 2.00 0 ATOM 3725 CB VAL 594 54,238 37,643 73,251 1.00 2.00 0 ATOM 3726 CG VAL 594 55,190 38,154 73,875 1.00 2.00 0 ATOM 3727 CG2 VAL 594 55,190 38,154 73,875 1.00 2.00 0 ATOM 3730 N ASP 595 56,112 38,991 71,356 1.00 2.00 0 ATOM 3730 N ASP 595 56,112 38,991 71,356 1.00 2.00 0 ATOM 3730 N ASP 595 56,108 37,919 71,356 1.00 3.06 0 ATOM 3730 C ASP 595 58,073 38,903 71,055 1.00 3.05 0 ATOM 3736 ODZ ASP 595 59,256 36,813 71,901 1.00 2.00 0 ATOM 3736 ODZ ASP 595 55,256 36,813 71,901 1.00 2.00 0 ATOM 3738 O ASP 595 55,355 38,913 37,907 72,601 1.00 2.00 0 ATOM 3741 CA ARG 596 57,342 41,927 71,460 1.00 2.00 0 ATOM 3741 CA ARG 596 57,342 41,927 71,460 1.00 2.00 0 ATOM 3743 CG ARG 596 55,468 43,096 70,716 1.00 2.00 0 ATOM 3741 CA ARG 596 55,468 43,096 70,716 1.00 2.00 0 ATOM 3742 CB ARG 596 55,468 43,096 70,716 1.00 2.00 0 ATOM 3745 CB ARG 596 55,468 43,096 70,716 1.00 2.00 0 ATOM 3745 CB ARG 596 55,468 43,096 70,716 1.00 2.00 0 ATOM 3745 CB ARG 596 55,468 43,096 70,716 1.0											-
XTOM 3716 CE2 TYR \$93 48,284 32,772 70,822 1.00 2.00 0 ATOM 3716 CE TYR \$93 48,856 31,540 71,013 1.00 2.00 0 ATOM 3720 C TYR \$93 \$1,955 30,0459 70,412 1.00 2.00 0 ATOM 3721 O TYR \$93 \$1,526 38,079 72,560 1.00 2.00 0 ATOM 3724 CA VAL \$94 \$1,213 37,463 73,420 1.00 2.00 0 ATOM 3726 CGI VAL \$94 \$1,313 36,599 74,501 1.00 30,066 0 ATOM 3728 C VAL \$94 \$1,333 36,599 74,501 1.00 30,066 0 ATOM 3732 C AL \$94 \$1,333 \$1,526 \$1,333 \$1,00									71.793	1.00 2.00	0
XTOM 3718 OR TYR 593	ATOM	3715	CD2		593						-
ATOM 3720 C TYR 593	-										-
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	MOTA	37 9 9	CD1	LEU	601		54.290	38.405	02.228	1.00 2.00	U



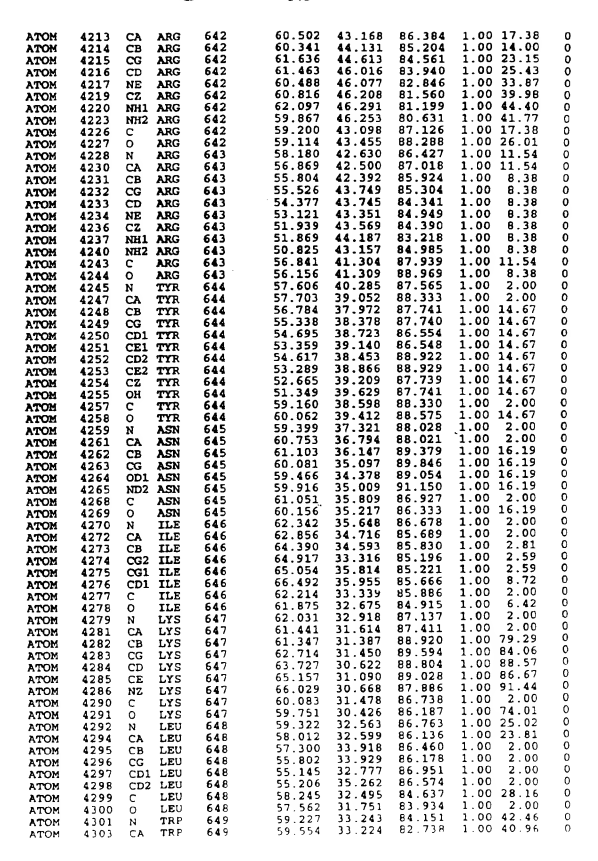


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					20 215	26 517	72.812	1 00	14.67	0
MOTA	3962	N	PHE	618 618	38. 2 15 39. 19 2	26.517 27.596	72.736		14.67	Ö
MOTA	3964	CA	PHE	618	39.146	28.387	74.044	1.00	2.00	ŏ
MOTA	3965 3966	CB CG	PHE	618	40.042	29.585	74.073	1.00	2.00	Ö
MOTA MOTA	3967		PHE	61B	39.501	30.871	74.058	1.00	2.00	Ō
ATOM	3968	CD2		618	41.422	29.441	74.147	1.00	2.00	0
ATOM	3969		PHE	618	40.322	32.007	74.119	1.00	2.00	0
ATOM	3970	CE2	PHE	618	42.254	30.562	74.209	1.00	2.00	0
MOTA	3971	CZ	PHE	618	41.696	31.854	74.195	1.00	2.00	0
ATOM	3972	Č	PHE	618	40.540	26.907	72.570	1.00	14.67	0
ATOM	3973	Ō	PHE	618	40.893	26.066	73.394	1.00	2.00	0
ATOM	3974	N	PHE	619	41.285	27.239	71.514	1.00	2.00	0
ATOM	3976	CA	PHE	619	42.583	26.599	71.271	1.00	2.00	0
MOTA	3977	CB	PHE	619	42.468	25.622	70.111	1.00	2.00	0
MOTA	3978	CG	PHE	619	41.404	24.595	70.311	1.00	2.00	0
MOTA	3 97 9	CD1		619	40.063	24.936	70.159	1.00	2.00	0
MOTA	3980		PHE	619	41.738	23.291	70.678 70.370	1.00	2.00 2.00	0
MOTA	3981	CE1		619	39.068	23.997 22.337	70.892	1.00	2.00	0
MOTA	3982	CE2	PHE	619	40.750 39.408	22.689	70.738	1.00	2.00	ŏ
MOTA	3983	CZ	PHE	619	43.719	27.566	71.005	1.00	2.00	ŏ
MOTA	3984	C	PHE	619	43.713	28.698	70.564	1.00	2.00	ŏ
MOTA	3985	0	PHE	619 620	44.936	27.115	71.282	1.00	2.00	ŏ
MOTA	3986	N	LEU	620	46.118	27.933	71.073	1.00	2.00	ŏ
MOTA	3988	CA	LEU	620	46.647	28.507	72.393	1.00	2.00	ŏ
MOTA	3989 3990	CB. CG	LEU	620	45.826	29.389	73.321	1.00	2.00	ŏ
MOTA MOTA	3991		LEU	620	46.741	29.868	74.437	1.00	2.00	Õ
ATOM	3992		LEU	620	45.247	30.563	72.565	1.00	2.00	0
ATOM	3993	C	LEU	620	47.225	27.093	70.438	1.00	2.00	0
MOTA	3994	ŏ	LEU	620	47.548	25.987	70.910	1.00	2.00	0
MOTA	3995	N	LEU	621	47.805	27.634	69.376	1.00	2.00	0
ATOM	3997	CA	LEU	621	48.880	26.971	68.669	1.00	2.00	0
MOTA	3998	CB	LEU	621	48.577	26.929	67.162	1.00	2.00	0
MOTA	3999	CG	LEU	621	47.256	26.280	66.739	1.00	2.00	0
MOTA	4000		LEU	621	47.021	26.511	65.253	1.00	2.00	0
MOTA	4001	CD2	LEU	621	47.281	24.813	67.062	1.00	2.00	0
MOTA	4002	С	LEU	621	50.162	27.756	68.943	1.00	2.00	0
MOTA	4003	0	LEU	621	50.121	28.937	69.285 68.774	1.00	2.00	ő
ATOM	4004	N	ARG	622	51.296 52.593	27.092	69.013	1.00	2.00	ŏ
MOTA	4006	CA	ARG	622 622	53.620	26.578	69.231	1.00	5.18	Ö
MOTA	4007	CB CC	ARG	622	54.967	27.052	69.716	1.00	5.18	ō
MOTA	4008	CG	ARG ARG	622	55.802	25.891	70.196	1.00	5.18	0
ATOM	4009 4010	CD NE	ARG	622	56.969	26.357	70.941	1.00	5.18	0
ATOM ATOM	4012	CZ	ARG	622	57.830	25.553	71.557	1.00	5.18	0
ATOM	4013	NH1	ARG	622	57.659	24.240	71.525	1.00	6.18	0
ATOM	4016	NH2	ARG	622	58.870	26.056	72.199	1.00	5.18	0
MOTA	4019	С	ARG	622	53. 0 70	28.615	67.894	i.ùO	2.00	0
ATOM	4020	ŏ	ARG	622	53.032	28.270	66.702	1.00	16.40	0
MOTA	4021	N	GLY	62 3	53.508	29.805	68.286	1.00	19.04	0
ATOM	4023	CA	GLY	623	54.043	30. 749	67.328	1.00	20.78	0
ATOM	4024	С	GLY	623	55.551	30.573	67.390		18.09	0
MOTA	4025	0	GLY	623	56. 06 9	29.805	68.212	1.00	2.00	0
ATOM	4026	N	asn	624	56.281	31.266	66.530	1.00	32.06	0
MOTA	4028	CA	ASN	624	57. 72 8	31.144	66.566		32.95	Ö
MOTA	4029	CB	asn	624	58.345	31.543	65.220		12.63	Ö
MOTA	4030	CG	ASN	624	58.120	33.002	64.857 65.538	1.00	8.84 9. 4 7	ő
MOTA	4031		ASN	624	57.381	33.754	63.757	1.00		Ö
MOTA	4032		ASN	624	58.721	33.411 31.988	67.721		32.42	ő
MOTA	4035	C	ASN	624	58.260 59.442	31.986	68.066	1.00	7.81	Ö
MOTA	4036	0	ASN	624 625	57.359	32.757	68.335	1.00	7.73	ő
MOTA	4037	N	HIS	625 625	57.687	33.614	69.477	1.00	8.83	Õ
MOTA	4039	CA	HIS HIS	625	57.030	34.987	69.320	1.00	2.00	Ō
ATOM	4040	CB CG	HIS	625	57.909	35.998	68.656	1.00	2.00	Ō
MOTA	4041 4042		HIS	625	59.042	35.854	67.933	1.00	2.00	0
ATOM ATOM	4042		HIS	625	57.668	37.351	68.721	1.00	2.00	0
ATOM	4045		HIS	625	58.618	37.999	68.069	1.00	2.00	0
F/ 1 O1/1	4040		,							

MOTA	4046	NE2	HIS	625	59.464	37.113	67.582	1.00 2.00	0
MOTA	4048	С	HIS	625	57.278	32.980	70.807 71.825	1.00 8.94 1.00 2.00	0
MOTA	4049	0	HIS	625	57.188 56.984	33.644 31.691	70.767	1.00 2.00	Ö
MOTA	4050	N	GLU	626	56.629	30.914	71.945	1.00 2.00	ŏ
MOTA	4052	CA	GLU	626 626	55.404	30.026	71.666	1.00 23.06	ŏ
MOTA	4053	CB	GLU	626	54.051	30.627	72.039	1.00 18.70	ŏ
MOTA	4054	CG	GLU	626	53.812	31.997	71.445	1.00 17.00	ō
MOTA	4055 4056	CD OE1	GLU	626	53.516	32.922	72.217	1.00 20.31	0
MOTA	4057	OE2	GLU	626	53.911	32.165	70.214	1.00 23.26	0
MOTA MOTA	4058	C	GLU	626	57.888	30.068	72.029	1.00 2.00	0
ATOM	4059	ŏ	GLU	626	57.851	28.841	71.921	1.00 23.50	0
MOTA	4060	N	CYS	627	59.015	30.74B	72.184	1.00 2.00	0
MOTA	4062	CA	CYS	627	60.290	30.067	72.218	1.00 2.00	0
MOTA	4063	CB	CYS	627	60.832	29.957	70.789	1.00 7.85 1.00 19.90	0
MOTA	4064	SG	CYS	627	62.235	28.843	70.575 73.105	1.00 19.90	Ö
MOTA	4065	Ċ	CYS	627	61.260 61.355	30.842 32.066	73.008	1.00 8.12	ŏ
MOTA	4066	0	CYS	627	61.971	30.116	73.970	1.00 19.49	ŏ
MOTA	4067	N	ALA	628 628	62.934	30.702	74.900	1.00 19.49	0
MOTA	4069	CA	ALA	628	63.704	29.601	75.567	1.00 2.00	0
MOTA	4070	CB C	ALA	628	63.890	31.662	74.206	1.00 19.49	0
ATOM ATOM	4071 4072	Ö	ALA	628	63.921	32.859	74.484	1.00 2.00	0
ATOM	4073	Ň	SER	629	64.660	31.101	73.291	1.00 10.83	0
ATOM	4075	CA	SER	629	65.637	31.813	72.478	1.00 14.60	0
ATOM	4076	СВ	SER	629	66.057	30.877	71.355	1.00 15.99	0
ATOM	4077	OG	SER	629	65.846	29.522	71.762	1.00 20.43 1.00 19.16	0
MOTA	4079	С	SER	629	65.119	33.131	71.899 71.759	1.00 19.16 1.00 13.43	ŏ
MOTA	4080	0	SER	629	65.869	34.085 33.167	71.755	1.00 2.00	ŏ
MOTA	4081	N	ILE	630	63.834 63.198	34.354	71.022	1.00 2.00	Ō
MOTA	4083	CA	ILE	630 630	62.076	33.973	70.017	1.00 2.00	0
MOTA	4084	CB	ILE ILE	630	61.675	35.174	69.202	1.00 2.00	0
MOTA	4085 4086	CG2 CG1	ILE	630	62.604	32.998	68.980	1.00 2.00	0
MOTA MOTA	4087	CD1		630	63.664	33.598	68. 08 0、	1.00 2.00	0
ATOM	4088	c	ILE	630	62.636	35.281	72.131	1.00 2.00	0
ATOM	4089	ō	ILE	630	62.8 6 6	36.492	72.068	1.00 2.00	0
MOTA	4090	N	ASN	631	61.932	34.729	73.137	1.00 10.29 1.00 9.61	ő
ATOM	4092	CA	asn	631	61.350	35.504	74.272 75.349	1.00 9.61 1.00 8.67	ŏ
MOTA	4093	CB	ASN	631	60.759	34.597 33.848	74.891	1.00 17.37	ō
ATOM	4094	CG	ASN	631	59. 55 5 59. 18 9	33.896	73.722	1.00 21.10	0
MOTA	4095		ASN	631	58.926	33.124	75.815	1.00 15.25	0
MOTA	4096		ASN	631 631	62.367	36.370	75.003	1.00 6.75	0
MOTA	4099	C	asn Asn	631	62.059	37.486	75.438	1.00 6.80	0
MOTA	4100 4101	и. О	ARG	632	63.555	35.810	75.189	1.00 2.00	0
MOTA MOTA	4103	CA	ARG	632	64.635	36.492	75.867	1.00 2.00	0
MOTA	4104	CB	ARG	632	65.873	35.595	75.909	1.00 6.06 1.00 8.20	0
MOTA	4105	CG	ARG	632	66.361	35.244	77.316	1.00 B.20 1.00 B.31	Ö
MOTA	4106	CD	ARG	632	67.436	36.202	77. 83 9 77. 91 8	1.00 11.75	ŏ
MOTA	4107	NE	ARG	632	67.003	37.598 38.635	77.907	1.00 13.36	0
MOTA	4109	CZ	ARG	632	67. 83 7 69. 144	38.423	77.811	1.00 13.50	0
MOTA	4110	NH1		632 632	67.372	39.883	78.001	1.00 18.51	0
ATOM	4113	NH2	ARG	632	64.962	37.781	75.148	1.00 2.00	0
MOTA MOTA	4116 4117	С 0	ARG	632	64.930	38.865	75.745	1.00 12.49	0
ATOM	4118	N	ILE	633	65.234	37.658	73.852	1.00 21.20	0
MOTA	4120	CA	ILE	633	65. 60 8	38.788	73.014	1.00 19.86	0
ATOM	4121	CB	ILE	633	66.085	38.308	71.610	1.00 32.61 1.00 29.80	ő
MOTA	4122	CG		633	66.132	39.471	70.625 71.700	1.00 28.84	Ö
ATOM	4123	CG:		633	67.490	37.716	72.594	1.00 35.23	ő
ATOM	4124	CD:		63 3	67. 619	36.496	72.815	1.00 22.19	ŏ
MOTA	4125	C	ILE	633	64. 55 0 64. 864	39.866 41.053	72.878	1.00 30.44	0
MOTA	4126	0	ILE	633	63.303	39.473	72.588	1.00 32.62	0
MOTA	4127	N	TYR	634 634	62.276	40.471	72.316	1.00 31.19	0
MOTA	4129	CA	TYR	634	61.395		71.147	1.00 13.98	0
MOTA	4130	CB	TYR TYR	634	62.231	39.755	69.941	1.00 19.07	0
MOTA	4131	CG	114	U J ¬	- -				

ATOM	4132		TYR	634	62.652	40.805	69.147	1.00 17.28 1.00 16.13	0
MOTA	4133	CE1	TYR	634	63.498 62.668	40.587 38.474	68.074 69.639	1.00 15.20	0
ATOM	4134	CD2	TYR	634 634	63.515	38.236	68.571	1.00 18.14	ō
MOTA	4135 4136	CE2	TYR TYR	634	63.935	39.296	67.786	1.00 21.41	0
ATOM ATOM	4137	OH	TYR	634	64.789	39.069	66.720	1.00 20.11	0
ATOM	4139	Ċ	TYR	634	61.430	41.025	73.446	1.00 28.63	0
ATOM	4140	ō	TYR	634	60.380	41.637	73.188	1.00 19.20	0
ATOM	4141	N	GLY	635	61.851	40.807	74.690	1.00 38.67	0
MOTA	4143	CA	GLY	635	61.114	41.407	75.786 76.780	1.00 39.39 1.00 41.31	0
ATOM	4144	Ç	GLY	635	60.270 60.103	40.658 41.155	77.892	1.00 69.69	Ö
MOTA	4145	0	GLY	635 636	59.725	39.502	76.431	1.00 54.49	Ö
ATOM	4146	и Су	PHE PHE	636	58.905	38.786	77.404	1.00 56.26	Ō
ATOM ATOM	4148 4149	CB	PHE	636	58.386	37.461	76.831	1.00 2.00	0
ATOM	4150	CG	PHE	636	57.275	36.845	77.639	1.00 2.00	0
ATOM	4151	CD1	PHE	636	56 .29 3	37.638	78.208	1.00 2.00	0
ATOM	4152	CD2	PHE	636	57.214	35.478	77.824	1.00 2.00	0
MOTA	4153	CE1	PHE	636	55.276	37.076	78.942	1.00 2.00	0
MOTA	4154	CE2	PHE	636	56.210	34.919 35.716	78.549 79.111	1.00 2.00 1.00 2.00	Ö
MOTA	4155	CZ	PHE	636	55.238 59.761	38.522	78.646	1.00 55.84	ŏ
ATOM	4156	C	PHE	636 636	59.321	38.728	79.791	1.00 2.00	ŏ
MOTA	4157 4158	O N	PHE TYR	637	61.005	38.114	78.415	1.00 2.00	0
MOTA MOTA	4160	CA	TYR	637	61.918	37.836	79.506	1.00 2.00	0
ATOM	4161	CB	TYR	637	63.266	37.419	78.947	1.00 18.03	0
ATOM	4162	CG	TYR	637	64.345	37.308	79.986	1.00 13.39	0
ATOM	4163	CD1	TYR	637	64.560	36.116	80.668	1.00 15.45	0
MOTA	4164	CE1	TYR	637	65.563	36.007 38.390	81.603 80.273	1.00 14.13 1.00 14.83	Ö
MOTA	4165	CD2	TYR	637	65.163 66.163	38.292	81.200	1.00 13.98	ŏ
MOTA	4166	CE2	TYR	637 637	66.363	37.101	81.863	1.00 14.88	ŏ
MOTA	4167 4168	CZ OH	TYR TYR	637	67.379	37.015	82.784	1.00 13.10	0
ATOM ATOM	4170	C	TYR	637	62.091	39.056	80.411	1.00 2.00	0
ATOM	4171	ŏ	TYR	637	62.362	38.920	81.605	1.00 20.03	0
ATOM	4172	N	ASP	638	61.919	40.240	79.833	1.00 2.00	0
MOTA	4174	CA	ASP	638	62.086	41.484	80.554	1.00 2.00 1.00 57.25	0
MOTA	4175	CB	ASP	638	62.701	42.522 42.053	79.614 79.046	1.00 66.80	ő
ATOM	4176	CG	ASP	638 638	64.049 64.078	41.539	77.905	1.00 65.06	ō
MOTA	4177	OD1	ASP ASP	638	65.077	42,175	79.750	1.00 69.12	0
MOTA	4178 4179	OD2 C	ASP	638	60.816	41.961	81.245	1.00 2.00	0
MOTA MOTA	4180	Ö	ASP	638	60.884	42.531	82.331	1.00 55.83	0
ATOM	4181	N	GLU	639	59.656	41.724	80.644	1.00 2.00	0
MOTA	4183	CA	GLU	639	58.405	42.074	81.317	1.00 2.00 1.00 64.74	0
MOTA	4184	CB	GLU	639	57.21 0 57.051	41.774 42.726	80. 41 9 79.261	1.00 64.74 1.00 71.70	ŏ
ATOM	4185	CG	GLU	639	55.900	42.720	78.355	1.00 66.55	ŏ
MOTA	4186	CD OE1	GLU	639 639	54.748	42.718	78.667	1.00 65.31	0
MOTA	4187 4188		GLU	639	56.152	41.684	77.329	1.00 72.66	0
MOTA MOTA	4189	C	GLU	639	58.372	41.145	82.547	1.00 2.00	0
ATOM	4190	õ	GLU	639	5B. 00 9	41.550	83.654	1.00 65.95	0
MOTA	4191	N	CYS	640	58.787	39.895	82.314	1.00 15.74 1.00 15.74	ő
MOTA	4193	CA	CYS	640	58.859	38.843 37.504	83.323 82.684	1.00 4.35	Ö
MOTA	4194	CB	CYS	640	59.187 57.734	36.587	82.208	1.00 13.44	0
MOTA	4195	SG	CYS CYS	640 640	59.859	39.081	84.426	1.00 15.74	0
MOTA	4196	C	CYS	640	59.590	38.723	85.564	1.00 11.73	0
MOTA MOTA			C 1 2		61.028	39.631	84.110	1.00 2.00	0
ATA WA'A	4197 4198		LYS	641	02.020				
	4198	N	LYS LYS	641 641	62.009	39.890	85.153	1.00 2.00	0
MOTA				641 641	62.009 63. 42 5	39.890 39.454	84.721	1.00 32.33	0
	4198 4200 4201 4202	N CA	LYS LYS LYS	641 641 641	62.009 63.425 64.225	39.890 39.454 40.436	84.721 83.869	1.00 32.33 1.00 32.55	0 0
ATOM ATOM ATOM ATOM	4198 4200 4201 4202 4203	N CA CB CG CD	LYS LYS LYS	641 641 641	62.009 63.425 64.225 65.735	39.890 39.454 40.436 40.154	84.721 83.869 83.954	1.00 32.33 1.00 32.55 1.00 38.99	0 0 0
MOTA MOTA MOTA MOTA MOTA	4198 4200 4201 4202 4203 4204	N CA CB CG CD CE	LYS LYS LYS LYS	641 641 641 641	62.009 63.425 64.225 65.735 66.281	39.890 39.454 40.436 40.154 40.198	84.721 83.869 83.954 85.386	1.00 32.33 1.00 32.55 1.00 38.99 1.00 41.14	0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	4198 4200 4201 4202 4203 4204 4205	N CA CB CG CD CE NZ	LYS LYS LYS LYS LYS	641 641 641 641 641	62.009 63.425 64.225 65.735 66.281 66.228	39.890 39.454 40.436 40.154 40.198 41.554	84.721 83.869 83.954 85.386 86.007	1.00 32.33 1.00 32.55 1.00 38.99	0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4198 4200 4201 4202 4203 4204 4205 4209	N CB CG CD CE NZ C	LYS LYS LYS LYS LYS LYS	641 641 641 641 641 641	62.009 63.425 64.225 65.735 66.281 66.228 61.950	39.890 39.454 40.436 40.154 40.198 41.554 41.371	84.721 83.869 83.954 85.386	1.00 32.33 1.00 32.55 1.00 38.99 1.00 41.14 1.00 38.23	0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	4198 4200 4201 4202 4203 4204 4205	N CA CB CG CD CE NZ	LYS LYS LYS LYS LYS	641 641 641 641 641	62.009 63.425 64.225 65.735 66.281 66.228	39.890 39.454 40.436 40.154 40.198 41.554	84.721 83.869 83.954 85.386 86.007 85.568	1.00 32.33 1.00 32.55 1.00 38.99 1.00 41.14 1.00 38.23 1.00 2.00	0 0 0 0 0 0

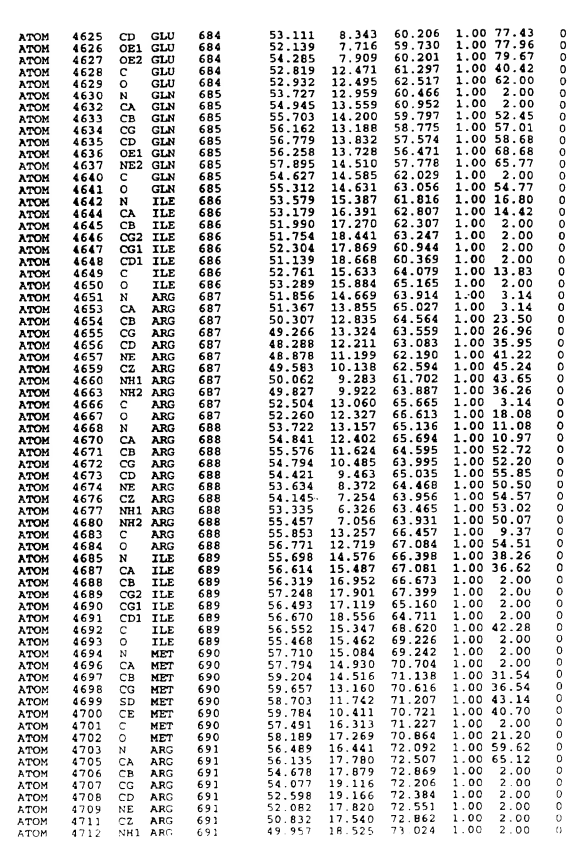


MOTA MOTA MOTA	4304 4305 4306	CB CG CD2	TRP TRP TRP	649 649 649	60.719 61.206 62.393	34.161 34.051 33.394	82.465 81.081 80.653	1.00 11.77 1.00 14.98 1.00 14.05 1.00 12.39	0 0
ATOM	4307	CE2	TRP	649	62.460	33.527	79.246 81.320	1.00 12.39 1.00 25.06	Ö
MOTA	4308		TRP	649	63. 41 2 60.607	32.703 34.546	79.951	1.00 17.81	Ō
MOTA	4309	CD1 NE1	TRP	649 649	61.356	34.232	78.846	1.00 15.30	0
ATOM ATOM	4310 4312	CZ2	TRP	649	63.510	32.995	78.495	1.00 13.58	0
ATOM	4313	CZ3	TRP	649	64.456	32.174	80.576 79.175	1.00 15.93 1.00 16.10	0
ATOM	4314	CH2	TRP	649	64.497 59.900	32.323 31.802	82.292	1.00 40.42	ŏ
MOTA	4315	C	TRP TRP	6 4 9 6 4 9	59.392	31.315	81.285	1.00 12.84	0
MOTA MOTA	4316 4317	N	LYS	650	60.748	31.138	83.069	1.00 2.00	0
ATON	4319	CA	LYS	650	61.179	29.776	82.796	1.00 2.00 1.00 19.18	0
ATOM	4320	CB	LYS	650	62.257	29.371 30.343	83.795 83.820	1.00 19.18	ŏ
MOTA	4321	CG	LYS	650 650	63.420 64.541	29.898	84.718	1.00 3.83	Ö
MOTA	4322 4323	CD CE	LYS LYS	650	65.148	28.593	84.221	1.00 11.09	0
ATOM ATOM	4324	NZ	LYS	650	66.096	27.974	85.202	1.00 11.03	0
ATOM	4328	C	LYS	650	59.976	28.851	82.892	1.00 2.00 1.00 5.16	0
MOTA	4329	0	LYS	650	59.926	27.805 29.234	82.239 83.702	1.00 10.53	ő
MOTA	4330	N	THR	651 651	58.997 57.792	28.422	83.838	1.00 14.10	Ö
MOTA	4332 4333	CA CB	THR THR	651	56.915	28.882	85.038	1.00 23.55	0
MOTA MOTA	4334	OG1		651	57.613	28.622	86.268	1.00 23.26	0
ATOM	4336	CG2		651	55.582	28.150	85.047 82.539	1.00 24.53 1.00 14.03	Ö
ATOM	4337	C	THR	651	57.012 56.599	28.543 27.534	81.951	1.00 23.01	ŏ
MOTA	4338	O N	THR PHE	651 652	56.837	29.779	82.082	1.00 2.00	0
MOTA MOTA	4339 4341	CA	PHE	652	56.127	30.032	80.840	1.00 2.00	0
MOTA	4342	CB	PHE	652	56.225	31.511	80.440 80.821	1.00 2.00 1.00 2.00	0
MOTA	4343	CG	PHE	652	55.027 55.164	32.320 33.486	81.536	1.00 2.00	ŏ
MOTA	4344		PHE	652 652	53.753	31.919	80.459	1.00 2.00	0
ATOM ATOM	4345 4346	CD2 CE1		652	54.039	34.245	81.884	1.00 2.00	0
ATOM	4347	CE2		652	52.630	32.678	80.808	1.00 2.00 1.00 2.00	0
MOTA	4348	CZ	PHE	652	52.775	33.832 29.160	81.515 79.743	1.00 2.00 1.00 2.00	ŏ
MOTA	4349	C	PHE	652 652	56.71 7 55. 9 81	28.456	79.067	1.00 2.00	0
MOTA	4350 4351	N O	PHE THR	653	58.039		79.608	1.00 2.00	0
MOTA MOTA	4353	CA	THR	653	58.631	28.344	78.552	1.00 2.00	0
MOTA	4354	CB	THR	653	60.126	28.538	78.429 79.255	1.00 2.14 1.00 2.14	ő
MOTA	4355	OG1		653	60.804 60.499	27.592 29.945	78.787	1.00 2.14	0
MOTA	4357 4358	CG2 C	THR	653 653	58.371	26.846	78.588	1.00 2.00	0
MOTA MOTA	4359	Ö	THR	653	58.452	26.196	77.555	1.00 4.20	0
ATOM	4360	N	ASP	654	58.064	26.271	79.743 79.749	1.00 2.00 1.00 2.00	Ö
MOTA	4362	CA	ASP	654	57.803 58.083	24.839 24.225	81.119	1.00 25.83	Ō
MOTA	4363	CB CG	ASP ASP	654 654	58.607	22.794	81.018	1.00 26.00	0
MOTA MOTA	4364 4365		ASP	654	59.163	22.420	79.960	1.00 29.22	0
ATOM	4366		ASP	654	58.472	22.043	82.006 79.332	1.00 28.12 1.00 2.00	Ö
MOTA	4367	C	ASP	654	56.367 56.014	24.603 23.518	78.874	1.00 18.75	0
MOTA	4368	0	ASP CYS	654 655	55.537	25.625	79.503	1.00 27.58	0
MOTA MOTA	4369 4371	N CA	CYS	655	54.146	25.546	79.095	1.00 27.58	0
MOTA	4372	СВ	CYS	655	53.333	26.675	79.722	1.00 8.45 1.00 8.45	0
MOTA	4373	SG	CYS	655	51.756	26.978 25.683	78.901 77.575	1.00.27.58	Ŏ
MOTA	4374	Ç	CYS	655 655	54.162 53.565	24.863	76.865	1.00 8.45	0
MOTA	4375 4376	O N	CYS PHE	655 656	54.871	26.709	77.088	1.00 7.64	0
MOTA MOTA	4378	CA	PHE	656	55.018	26.966	75.653	1.00 7.64 1.00 12.44	0
MOTA	4379	CB	PHE	656	55.967	28.145	75.402 75.747	1.00 12.44	0
MOTA	4380	CG	PHE	656	55.384 54.073	29.483 29.591	76.242	1.00 12.44	Ö
MOTA	4381	CD:		656 656	56.144	30.641	75.594	1.00 12.44	0
MOTA MOTA	4382 4383	CE:		656	53.525	30.840	76.585	1.00 12.44	0
MOTA	4384	CE		656	55.614	31.891	75.929	1.00 12.44	U

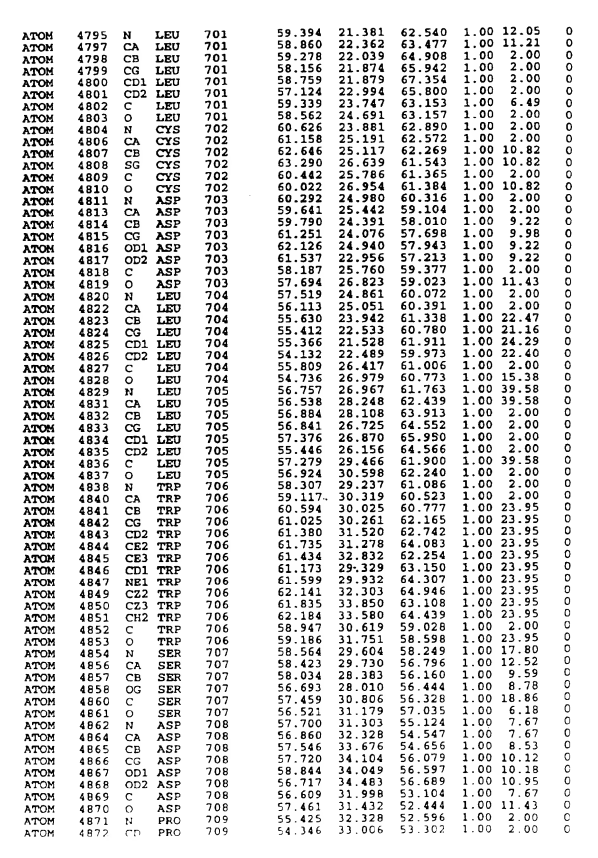
ATOM	4385	CZ	PHE	656	54.296	31.990	76.430	1.00 12.44	0
ATOM	4386	C	PHE	656	55.564	25.705	74.961	1.00 7.64	0
MOTA	4387	0	PHE	656	55.033	25.274	73.940	1.00 12.44	0
MOTA	4388	N	ASN	657	56.595	25.099	75.543	1.00 2.00	0
MOTA	4390	CA	ASN	657	57.198	23.896	75.006	1.00 2.00	0
MOTA	4391	CB	asn	657	58.353	23.425	75.892	1.00 12.64	0
MOTA	4392	CG	ASN	657	59.614	24.233	75.690	1.00 12.64	0
MOTA	4393	OD1	asn	657	59.564	25.404	75.350	1.00 12.64	0
MOTA	4394	ND2	asn	657	60.757	23.607	75.896	1.00 12.64	0
ATOM	4397	С	asn	657	56.202	22.772	74.881	1.00 2.00	0
ATOM	4398	0	asn	657	56.585	21.668	74.532	1.00 12.64	0
MOTA	4399	N	CYS	658	54.934	23.017	75.196	1.00 2.00	0
MOTA	4401	CA	CYS	658	53.930	21.964	75.080	1.00 2.00	0
MOTA	4402	CB	CYS	658	53.543	21.46B	76.475	1.00 10.87	0
MOTA	4403	SG	CYS	658	55.004	20.919	77.397	1.00 10.87	0
MOTA	4404	C	CYS	658	52.708	22.390	74.258	1.00 2.00	0
MOTA	4405	0	CYS	658	51.726	21.659	74.162	1.00 10.87	0
MOTA	4406	N	LEU	659	52.799	23.560	73.634	1.00 11.17	0
ATOM	4408	CA	LEU	659	51.739	24.085	72.781	1.00 11.17	0
MOTA	4409	CB	LEU	659	51.998	25.569	72.470	1.00 2.00	0
ATOM	4410	CG	LEU	659	51.757	26.629	73.542 73.123	1.00 2.00	0
ATOM	4411	_	LEU	659	52.286	27.965		1.00 2.00	0
MOTA	4412	CD2	LEU	659	50.307	26.759	73.767	1.00 2.00	0
MOTA	4413	Ċ	LEU	659	51.651	23.298	71.450	1.00 11.17	0
ATOM	4414	0	LEU	659	52.681	22.845	70.900 70.931	1.00 2.00 1.00 14.96	0
ATOM	4415	N	PRO	660	50.415	23.109	70.531	1.00 14.90	Ö
ATOM	4416	CD	PRO	660	49.134	23.535 22.398		1.00 2.00	Ö
MOTA	4417	CA	PRO	660	50.166 48.640	22.381	69.680 69.590	1.00 14.90	ŏ
ATOM	4418	CB	PRO	660	48.194	22.472	71.028	1.00 2.00	ŏ
MOTA	4419	CG	PRO	660	50.796	23.248	68.579	1.00 14.96	ŏ
MOTA	4420	C	PRO	660	50.736	24.474	68.709	1.00 2.00	ŏ
MOTA	4421	0	PRO	660 661	51.195	22.603	67.488	1.00 28.66	ŏ
MOTA	4422	N	ILE		51.880	23.279	66.397	1.00 30.55	ŏ
MOTA	4424	CA	ILE	661 661	53.126	22.480	66.036	1.00 19.83	ŏ
MOTA	4425	CB	ILE	661	54.064	22.432	67.228	1.00 22.82	ŏ
MOTA	4426	CG2	ILE	661	52.728	21.050	65.669	1.00 24.89	Õ
MOTA	4427 4428	CG1	ILE	661	53.896	20.161	65.299	1.00 29.65	Õ
MOTA MOTA	4429	CDI	ILE	661	51.074	23.577	65.132	1.00 29.24	0
MOTA	4430	Ö	ILE	661	51.428,	24.493	64.372	1.00 20.63	0
ATOM	4431	N	ALA	662	50.004	22.810	64.915	1.00 22.79	0
ATOM	4433	CA	ALA	662	49.112	22.970	63.751	1.00 22.79	0
ATOM	4434	CB	ALA	662	49.652	22.187	62.529	1.00 2.00	0
MOTA	4435	c	ALA	662	47.683	22.507	64.086	1.00 22.79	0
ATOM	4436	ō	ALA	662	47.439	21.887	65.139	1.00 2.00	0
ATOM	4437	N	ALA	663	46.739	22.816	63.205	1.00 2.00	0
MOTA	4439	CA	ALA	663	45.347	22.438	63.410	1.00 2.00	O
MOTA	4440	CB	ALA	663	44.599	23.535	64.124	1.00 18.31	0
MOTA	4441	C	ALA	663	44.748	22.223	62.053	1.00 2.00	0
MOTA	4442	0	ALA	663	45.323	22.623	61.041	1.00 22.14	0
MOTA	4443	N	ILE	664	43.600	21.565	62.028	1.00 16.69	0
MOTA	4445	CA	ILE	664	42.892	21.300	60.788	1.00 16.69	0
ATOM	4446	CB	ILE	664	43.240	19.885	60.208	1.00 9.81	0
ATOM	4447	CG2	ILE	664	42.396	19.597	58.976	1.00 9.81	0
ATOM	4448	CG1	ILE	664	44.724	19.810	59.819	1.00 9.81	0
MOTA	4449	CD1	ILE	664	45.161	18.450	59.316	1.00 9.81	0
ATOM	4450	С	ILE	664	41.408	21.397	61.118	1.00 16.69	0
MOTA	4451	0	ILE	664	40.880	20.603	61.899	1.00 9.81	0
MOTA	4452	N	VAL	665	40.754	22.408	60.561	1.00 14.09	0
ATOM	4454	CA	VAL	665	39.332	22.608	60.777	1.00 14.09	0
MOTA	4455	CB	VAL	665	38.954	24.091	60.742	1.00 2.00	0
ATOM	4456		VAL	665	37.450	24.239	60.710	1.00 2.00	0
ATOM	4457	CG2	VAL	665	39.500	24.788	61.950	1.00 2.00 1.00 14.09	0
MOTA	4458	C	VAL	665	38.509	21.873	59.729 58.524	1.00 14.09	. 0
ATOM	4459	0	VAL	665	38.768	21.972 21.137	60.219	1.00 2.00	0
MOTA	4460	N	ASP	666	37.517	21.137	59.407	1.00 2.00	Ö
MOTA	4462	CA	ASP	666	36.595	20.345	58.933	1.00 2.00	0
MOTA	4463	CB	ASP	6 66	35.443	41.433	ر د ر	1.00 00.04	U

MOTA	4464	CG	ASP	666		34.545	21.675	60.081		73.26	0
MOTA	4465	OD1		666		33.763	20.838	60.578		71.65	0
ATOM	4466	OD2	ASP	666		34.623 37.234	22.851 19.577	60. 49 3 58.235	1.00	75.52 2.00	0
MOTA	4467	C O	ASP ASP	666 666		36.648	19.467	57.158		57.66	0
MOTA MOTA	4468 4469	N	GLU	667		38.439	19.048	58.472		17.20	Ö
ATOM	4471	CA	GLU	667		39.203	18.270	57.489		17.69	ŏ
ATOM	4472	CB	GLU	667		38.455	16.981	57.138		42.87	0
MOTA	4473	CG	GLU	667		38.170	16.101	58.345		52.20	0
MOTA	4474	CD	GLU	667		37.457 36.222	14.806 14.717	57.988 58.211		53.85 51.15	0
MOTA MOTA	4475 4476	OE1 OE2	GLU	6 6 7 6 6 7		38.139	13.877	57.494		55.67	Ö
MOTA	4477	C	GLU	667		39.584	19.015	56.207		17.24	ŏ
MOTA	4478	ō	GLU	667		40.146	18.421	55.286		34.30	0
MOTA	4479	N	LYS	668		39.314	20.319	56.172		26.53	0
MOTA	4481	CA	LYS	668		39.615	21.133 21.709	55.002 54.410		20.22 13.48	0
MOTA	4482 4483	CB CG	LYS LYS	668 668		38.318 37.383	20.628	53.859		13.48	0
MOTA MOTA	4484	CD	LYS	668		38.119	19.779	52.819		13.48	ŏ
MOTA	4485	CE	LYS	668		37.341	18.551	52.398	1.00	16.60	Ō
MOTA	4486	NZ	LYS	668		38.247	17.564	51.739		18.66	0
MOTA	4490	Ç	LYS	668		40.636	22.244	55.240		19.61	0
MOTA	4491	0	LYS ILE	668 669		41.676 40.356	22.264 23.166	54.584 56.158	1.00	13.48	0
MOTA MOTA	4492 4494	N CA	ILE	669		41.295	24.263	56.424	1.00	2.00	ő
ATOM	4495	CB	ILE	669		40.617	25.456	57.183	1.00	2.00	ŏ
MOTA	4496	CG2	ILE	669		41.521	26.680	57.166	1.00	2.00	0
MOTA	4497	CG1	ILE	669		39.298	25.839	56.518	1.00	2.00	0
MOTA	4498	CD1	ILE	669		38.581 42.439	26.960 23.721	57.189 57.279	1.00	2.00	0
ATOM ATOM	4499 4500	C	ILE	669 6 6 9		42.201	23.034	58.269	1.00	2.00	ŏ
MOTA	4501	N	PHE	670	•	43.673	24.010	56.892	1.00	2.00	Ŏ
MOTA	4503	CA	PHE	670		44.841	23.551	57.641	1.00	2.00	0
MOTA	4504	CB	PHE	670		45.804	22.801	56.718	1.00	2.00	0
MOTA	4505	CG	PHE	670		47.182	22.614	57.291 58.007	1.00 1.00	2.00 2.00	0
MOTA	4506		PHE	670 670		47.503 48.163	21.473 23.569	57.093	1.00	2.00	ő
MOTA MOTA	4507 4508	CD2 CE1	PHE PHE	670		48.769	21.288	58.506	1.00	2.00	ŏ
ATOM	4509	CE2	PHE	670		49.436	23.384	57.596	1.00	2.00	0
MOTA	4510	CZ	PHE	670		49.737	22.241	58.302	1.00	2.00	0
MOTA	4511	С	PHE	67 0		45.528	24.780	58.190	1.00	2.00 2.00	0
ATOM	4512	0	PHE	670 671		45.901 45.725	25.652 24.839	57.421 59.501	1.00 1.00	2.00	ő
ATOM ATOM	451 3 451 5	N CA	CYS CYS	671		46.353	25.992	60.120	1.00	2.00	ō
MOTA	4516	CB	CYS	671		45.395	26.596	61.136		14.45	0
MOTA	4517	5G	CYS	671		43.708	26.779	60.568	1.00	25.33	Ó
MOTA	4518	C	CYS	671		47.685	25.701	60.816 61.310	1.00 1.00	2.00 8.01	0
MOTA	4519	0	CYS CYS	671 672		47.921 48.546	24.589 26.714	60.845	1.00	2.00	ő
ATOM ATOM	4520 4522	N C A	CYS	672			26.666	61.515	1.00	2.00	ŏ
ATOM	4523	CB	CYS	672		50.826	25.741	60.792	1.00	9.00	0
MOTA	4524	SG	CYS	672		51.494	26.372	59.265	1.00	11.24	0
MOTA	4525	C	CYS	672		50.336	28.122	61.525	1.00	2.00	0
MOTA	4526	0	CYS	6 7 2		49.751 51.369	28.963 28.440	60.850 62.300	1.00 1.00	9.00 17.94	0
ATOM ATOM	4527 4529	N CA	HIS HIS	6 7 3 673		51.855	29.817	62.360		17.94	ŏ
ATOM	4530	C	HIS	67 3		52.456	30.359	61.065	1.00	17.94	0
MOTA	4531	Ō	HIS	673		51.960	31.349	60.514	1.00	2.00	0
MOTA	4532	CB	HIS	67 3		52.894	29.965	63.459	1.00	2.00	0
MOTA	4533	CG	HIS	673		53.283	31.383 32.377	63.724 64.033	1.00	2.00 2.00	0
ATOM	4534 4536		HIS HIS	67 3 67 3		52.388 54.503	32.377	63.718	1.00	2.00	0
ATOM ATOM	4536 4537		HIS	6 7 3		54.371	33.330	64.019	1.00	2.00	Ö
ATOM	4538		HIS	673		53.072	33.512	64.199	1.00	2.00	0
ATOM	4539	N	GLY	674		53.545	29.721	60.626	1.00	2.00	0
ATOM	4541	CA	GLY	674		54.260	30.101	59.417	1.00	2.00	0
ATOM	4542	C	GLY	674 674		53.773 53.410	29.407 30.063	58.163 57.203	1.00	2.00 11.62	0
ATOM	4543	O	GLY	674		410.410	.000) I . LUJ	1.00	02	U

MOTA	4544	N	GLY	675	53.759	28.087	58.136	1.00 6.25	0
ATOM	4546	CA	GLY	675	53.286	27.440	56.931	1.00 6.25	Ŏ
ATOM	4547	C	GLY	675	53.823	26.061	56.604	1.00 6.25	Ŏ
ATOM	4548	õ	GLY	675	53.867	25.173	57.452	1.00 28.15	Ō
MOTA	4549	N	LEU	676	54.241	25.883	55.356	1.00 2.00	Ŏ
MOTA	4551	CA	LEU	676	54.723	24.591	54.886	1.00 2.00	Ō
ATOM	4552	CB	LEU	676	54.336	24.409	53.418	1.00 2.00	ŏ
ATOM	4553	CG	LEU	676	52.827	24.625	53.244	1.00 2.00	ō
ATOM	4554		LEU	676	52.412	24.556	51.777	1.00 2.00	ō
MOTA	4555	CD2	LEU	676	52.100	23.575	54.075	1.00 2.00	Õ
MOTA	4556	C	LEU	676	56.207	24.333	55.095	1.00 2.00	Ŏ
ATOM	4557	õ	LEU	676	56.981	25.248	55.381	1.00 2.00	Ō
ATOM	4558	N	SER	677	56.582	23.069	54.945	1.00 12.63	0
ATOM	4560	CA	SER	677	57.946	22.617	55.141	1.00 12.63	0
MOTA	4561	CB	SER	67 7	58.083	21.985	56.539	1.00 2.43	0
MOTA	4562	OG	SER	677	59.248	21.192	56.649	1.00 2.03	0
MOTA	4564	С	SER	677	58.247	21.556	54.095	1.00 12.63	0
ATOM	4565	0	SER	677	57.405	20.684	53.838	1.00 10.72	0
ATOM	4566	N	PRO	678	59.449	21.605	53.484	1.00 2.00	0
MOTA	4567	CD	PRO	678	60.474	22.638	53.681	1.00 15.59	0
MOTA	4568	CA	PRO	678	59.887	20.647	52.469	1.00 2.00	0
MOTA	4569	CB	PRO	678	61.329	21.076	52.184	1.00 15.59	0
ATOM	4570	CG	PRO	678	61.299	22.517	52.416	1.00 15.59	0
ATOM	4571	C	PRO	678	59.859	19.234	53.032	1.00 2.00	0
MOTA	4572	0	PRO	678	59.990	18.269	52.293	1.00 15.59	0
MOTA	4573	N	ASP	679	5 9.68 5	19.112	54.342	1.00 2.00	0
MOTA	4575	CA	ASP	679	59.687	17.813	54.992	1.00 2.00	0
MOTA	4576	CB	ASP	679	60.510	17.926	56.271	1.00 26.28	0
MOTA	4577	CG	ASP	679	61.766	18.758	56.073	1.00 24.05	0
MOTA	4578	OD1	ASP	679	62.687	18.269	55.383	1.00 32.92	0
MOTA	4579	OD2	ASP	679	61.826	19.902	56.586	1.00 28.53	0
MOTA	4580	C	ASP	679	58.303	17.278	55.314	1.00 2.00	0
MOTA	4581	0	ASP	679	58.129	16.094	55.576 55.279	1.00 24.04 1.00 2.00	Ö
MOTA	4582	N	LEU	680	57.315	18.149 17.747	55.622	1.00 2.00	ő
ATOM	4584	CA	LEU	680	55.970 55.147	19.000	55.916	1.00 2.00	ő
MOTA	4585	CB	LEU	680	53.683	18.788	56.262	1.00 2.86	ŏ
MOTA	4586	CG	LEU	680 680	53.540	17.791	57.398	1.00 2.86	ŏ
ATOM ATOM	4587 4588	CD1 CD2	LEU	680	53.085	20.134	56.582	1.00 2.86	ō
ATOM	4589	CDZ	LEU	680	55.262	16.862	54.588	1.00 2.00	Ō
ATOM	4590	ŏ	LEU	680	54.421	17.331	53.814	1.00 15.95	0
ATOM	4591	Ň	GLN	681	55.588	15.578	54.561	1.00 2.00	0
ATOM	4593	CA	GLN	681	54.922	14.686	53.611	1.00 2.00	0
ATOM	4594	CB	GLN	681	55.719	13.410	53.385	1.00 36.99	0
ATOM	4595	ĊĠ	GLN	681	57.097	13.620	52.863	1.00 36.99	0
ATOM	4596	CD	GLN	681	57.716	12.321	52.468	1.00 36.99	0
ATOM	4597	OE1	GLN	681	57.589	11.894	51.326	1.00 36.99	0
ATOM	4598	NE2	GLN	681	58.377	11.663	53.409	1.00 36.99	0
ATOM	4601	С	GLN	681	53.551	14.319	54.159	1.00 2.00	0
MOTA	4602	0	GLN	681	52.547	14.392	53.445	1.00 11.55	0
MOTA	4603	И	SER	682	53.505	13.923	55.423	1.00 39.00	0
MOTA	4605	CA	SER	682	52.238	13.566	56.027	1.00 42.03	0
MOTA	4606	CB	SER	682	52.131	12.048	56.224	1.00 2.00 1.00 2.00	Ö
MOTA	4607	OG	SER	682	53.198	11.507	56.993		ő
MOTA	4609	C	SER	682	52.003	14.284	57.343	1.00 38.75 1.00 2.00	ő
MOTA	4610	0	SER	682	52.941	14.742	58.009 57.6 9 8	1.00 15.16	Ö
MOTA	4611	N	MET	683	50.730	14.397	58.938	1.00 15.16	ŏ
ATOM	4613	CA	MET	683	50.338	15.029	59.044	1.00 11.68	Ö
MOTA	4614	CB	MET	683	48.820 48.099	15.069 15.565	57.799	1.00 12.75	ő
ATOM	4615	CG	MET	683	48.099	17.328	57.477	1.00 11.68	õ
MOTA	4616	SD	MET	683	49.486	17.429	56.200	1.00 12.61	ŏ
MOTA	4617	CE	MET	683 683	50.906	14.131	60.024	1.00 15.16	ŏ
MOTA	4618	C	MET	683	51.215	14.585	61.116	1.00 15.01	ŏ
ATOM	4619 4620	0	MET GLU	684	51.050	12.848	59.705	1.00 40.22	ŏ
ATOM ATOM	4622	N CA	GLU	684	51.597	11.881	60.644	1.00 41.85	ō
ATOM	4623	CB	GLU	684	52.007	10.594	59.937	1.00 63.04	0
MOTA	4624	CG	GLU	684	52.850	9.689	60.823	1.00 72.94	0



ATOM ATOM	4715 4718	С	ARG ARG ARG	691 691 691	50.482 56.930 57.176	16.273 18.632 19.794	73.046 73.458 73.100	1.00 2.00 1.00 64.88 1.00 2.00	0
MOTA	4719 4720	0	PRO	692	57.265	18.151	74.693	1.00 0.89	Ö
MOTA MOTA	4721	N CD	PRO	692	56.930	16.947	75.482	1.00 19.88	ŏ
MOTA	4722	CA	PRO	692	58.060	19.115	75.500	1.00 0.77	Ö
ATOM	4723	CB	PRO	692	58.305	18.357	76.811	1.00 21.87	ŏ
MOTA	4724	CG	PRO	692	57.071	17.462	76.911	1.00 19.02	ŏ
ATOM	4725	C	PRO	692	59.327	19.317	74.643	1.00 0.10	ŏ
MOTA	4726	Ö	PRO	692	60.258	18.502	74.690	1.00 22.15	ŏ
ATOM	4727	N	THR	693	59.310	20.362	73.811	1.00 2.00	Ō
ATOM	4729	CA	THR	693	60.395	20.605	72.896	1.00 2.00	0
ATOM	4730	CB	THR	693	60.153	19.842	71.569	1.00 39.91	0
ATOM	4731	OG1	THR	693	61.310	19.947	70.728	1.00 47.46	0
ATOM	4733	CG2	THR	693	58.944	20.419	70.830	1.00 46.31	0
MOTA	4734	С	THR	693	60.567	22.057	72.560	1.00 2.00	0
MOTA	4735	0	THR	693	59.640	22.848	72.678	1.00 41.07	0
ATOM	4736	N	ASP	694	61.782	22.399	72.154	1.00 4.51 1.00 4.51	0
ATOM	4738	CA	ASP	694	62.075 63.429	23.747 24.203	71.736 72.283	1.00 83.74	0
MOTA	4739	CB	ASP	694	63.337	25.520	73.041	1.00 83.74	ŏ
MOTA	4740	CG	ASP	694	63.231	25.484	74.285	1.00 83.74	ŏ
ATOM	4741		ASP	694 694	63.366	26.594	72.400	1.00 83.74	ŏ
ATOM	4742 4743	C C	ASP ASP	694	62.101	23.682	70.201	1.00 4.51	Ŏ
MOTA	4744	0	ASP	694	62.403	22.632	69.629	1.00 83.74	0
ATOM ATOM	4745	Ŋ	VAL	695	61.743	24.784	69.546	1.00 31.71	0
ATOM	4747	CA	VAL	695	61.760	24.864	68.087	1.00 37.86	0
ATOM	4748	CB	VAL	695	61.212	26.224	67.623	1.00 72.52	0
ATOM	4749		VAL	695	61.120	26.268	66.113	1.00 68.63	0
MOTA	4750	CG2	VAL	695	59.863	26.474	68.253	1.00 68.27	0
ATOM	4751	C	VAL	695	63.242	24.744	67.687	1.00 34.22	0
MOTA	4752	0	VAL	695	- 64.070	25.548	68.123	1.00 75.22 1.00 2.00	0
ATOM	4753	N	PRO	696	63.599	23.730	66.873 66.373	1.00 2.00 1.00 5.25	ŏ
MOTA	4754	CD	PRO	696	62.777	22.616 23.549	66.462		ŏ
ATOM	4755	CA	PRO	.696	64.998	22.163	65.803	1.00 5.25	ŏ
ATOM	4756	CB	PRO	696	64.997 63.768	21.486	66.381	1.00 5.25	Ō
ATOM	4757	CC	PRO	696 696	65.570	24.612	65.536	1.00 2.00	0
MOTA	4758	C O	PRO PRO	696	64.878	25.556	65.137	1.00 5.25	0
MOTA MOTA	4759 4760	N	ASP	697	66.85Q	24.431	65.214	1.00 34.70	0
ATOM	4762	CA	ASP	697	67.598	25.311	64.321	1.00 35.10	0
ATOM	4763	CB	ASP	697	69.098	24.951	64.376	1.00 81.36	0
ATOM	4764	CG	ASP	697	69.630	24.774	65.B25	1.00 81.80	0
MOTA	4765	OD1	ASP	697	69.612	23.613	66.347	1.00 0.89 1.00 0.05	ő
MOTA	4766	OD2	ASP	697	70.075	25.792	66.436 62.894	1.00 36.63	ŏ
MOTA	4767	C	ASP	697	67.037 67.014	25.099 26.022	62.069	1.00 0.75	ō
MOTA	4768	0	ASP	697	66.576	23.877	62.620	1.00 8.48	0
ATOM	4769	N	GLN	698	65.997	23.510	61.335	1.00 2.00	0
MOTA	4771	CA	GLN GLN	698 698	67.089	23.346	60.285	1.00 43.65	0
ATOM	4772 4773	CB CG	GLN	698	68.191	22.376	60.649	1.00 44.93	0
MOTA MOTA	4774	CD	GLN	698	69.158	22.169	59.501	1.00 43.25	0
MOTA				698	68.781	22.242	58.327	1.00 45.90	0
	4//5	OEI	מענט	0,0					
ATOM	4775 4776	NE2	GLN GLN	698	70.411	21.911	59.830	1.00 44.14	
MOTA MOTA	4776 4779			698 698	70. 41 1 65. 21 1	21.911 22.216	61.482	1.00 2.32	0
	4776	NE2	GLN	698 698 698	70.411 65.211 65.396	21.911 22.216 21.486	61.482 62.452	1.00 2.32 1.00 42.62	0
MOTA	4776 4779	NE2 C	GLN GLN GLY	698 698 698 699	70.411 65.211 65.396 64.324	21.911 22.216 21.486 21.939	61.482 62.452 60.530	1.00 2.32 1.00 42.62 1.00 2.00	0 0
ATOM ATOM	4776 4779 4780	NE2 C O N CA	GLN GLN GLY GLY	698 698 698 699	70.411 65.211 65.396 64.324 63.510	21.911 22.216 21.486 21.939 20.728	61.482 62.452 60.530 60.576	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00	0 0 0
MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784	NE2 C O N CA C	GLN GLN GLY GLY GLY	698 698 698 699 699	70.411 65.211 65.396 64.324 63.510 62.046	21.911 22.216 21.486 21.939 20.728 21.059	61.482 62.452 60.530 60.576 60.346	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00	0 0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785	NE2 C O N CA C	GLN GLN GLY GLY GLY GLY	698 698 698 699 699 699	70.411 65.211 65.396 64.324 63.510 62.046 61.726	21.911 22.216 21.486 21.939 20.728 21.059 22.204	61.482 62.452 60.530 60.576 60.346 60.013	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	0 0 0
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785 4786	NE2 C O N CA C O N	GLN GLN GLY GLY GLY GLY LEU	698 698 698 699 699 699 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.153	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090	61.482 62.452 60.530 60.576 60.346 60.013 60.545	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	00000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785 4786 4788	NE2 C O N CA C O N CA	GLN GLN GLY GLY GLY LEU LEU	698 698 699 699 699 699 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.153 59.700	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090 20.297	61.482 62.452 60.530 60.576 60.346 60.013 60.545 60.352	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	00000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785 4786 4788 4789	NE2 C O N CA C O N CA CB	GLN GLN GLY GLY GLY LEU LEU LEU	698 698 699 699 699 699 700 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.153 59.700 58.941	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090 20.297 18.962	61.482 62.452 60.530 60.576 60.346 60.013 60.545	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	0000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4776 4779 4780 4781 4783 4784 4785 4786 4788 4789	NE2 C O N CA C O N CA CB CB	GLN GLN GLY GLY GLY LEU LEU LEU	698 698 699 699 699 700 700 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.153 59.700 58.941 57.436	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090 20.297 18.962 18.836	61.482 62.452 60.530 60.576 60.346 60.013 60.545 60.352 60.555	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 4.64 1.00 8.84 1.00 8.28	0000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785 4786 4788 4789 4790 4791	NE2 C O N CA C O N CA CB CG CDI	GLN GLN GLY GLY GLY LEU LEU LEU LEU	698 698 699 699 699 700 700 700 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.153 59.700 58.941	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090 20.297 18.962	61.482 62.452 60.530 60.576 60.346 60.013 60.555 60.273 58.793 60.803	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 4.64 1.00 8.84 1.00 8.28 1.00 5.23	000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785 4786 4788 4790 4791 4792	NE2 C O N CA C O N CA CB CG CD1	GLN GLN GLY GLY GLY LEU LEU LEU LEU LEU	698 698 699 699 699 700 700 700 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.753 59.700 58.941 57.436 57.219	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090 20.297 18.836 18.836 18.881 17.536 21.400	61.482 62.452 60.530 60.576 60.346 60.013 60.545 60.352 60.273 58.793 60.803 61.247	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 8.84 1.00 8.28 1.00 5.23 1.00 2.00	0000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	4776 4779 4780 4781 4783 4784 4785 4786 4788 4789 4790 4791	NE2 C O N CA C O N CA CB CG CDI	GLN GLN GLY GLY GLY LEU LEU LEU LEU	698 698 699 699 699 700 700 700 700	70.411 65.211 65.396 64.324 63.510 62.046 61.726 61.153 59.726 57.436 57.219 56.879	21.911 22.216 21.486 21.939 20.728 21.059 22.204 20.090 20.297 18.962 18.836 18.881 17.536	61.482 62.452 60.530 60.576 60.346 60.013 60.555 60.273 58.793 60.803	1.00 2.32 1.00 42.62 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 4.64 1.00 8.84 1.00 8.28 1.00 5.23	000000000000

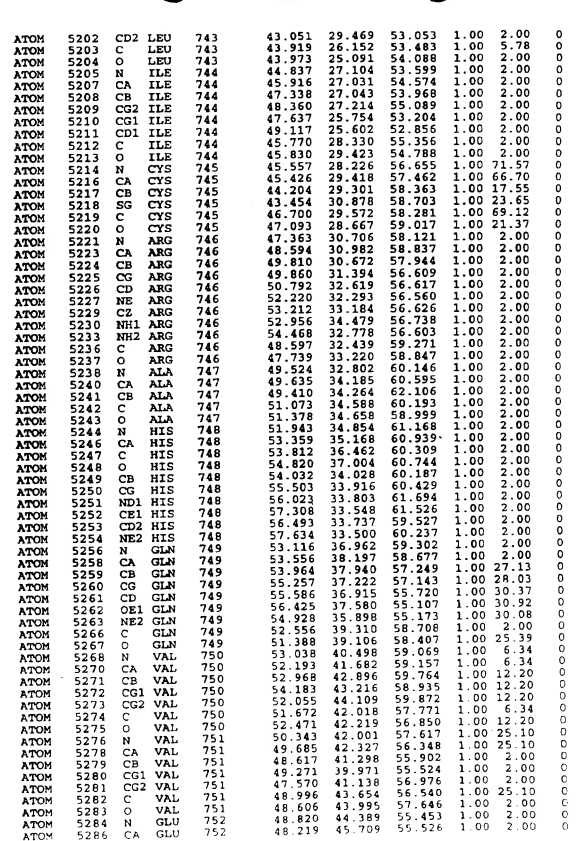


32.078 MOTA 4873 PRO 709 55.029 1.00 2.00 CA 51.214 53.507 32.036 51.283 2.00 MOTA 4874 CB PRO 709 1.00 4875 CG PRO 709 53.169 32.316 52.728 1.00 2.00 MOTA 1.00 33.269 50.406 MOTA 4876 С PRO 709 55.475 2.00 MOTA 4877 0 PRO 709 55.071 34.393 50.712 1.00 2.00 33.046 4878 710 56.306 49.393 1.00 2.57 ASP N MOTA 0 34.155 56.769 48.576 MOTA 4880 CA **ASP** 710 1.00 4.76 710 710 48.537 48.303 34.215 4881 CB ASP 58.297 1.00 18.70 MOTA 0 58.824 35.631 1.00 26.43 ASP MOTA 4882 CG 0 36.497 58.070 47.793 1.00 26.58 ATOM 4883 OD1 ASP 710 0 OD2 ASP 710 60.002 35.884 48.635 1.00 30.90 MOTA 4884 n 56.211 47.168 710 34.045 1.00 ATOM 4885 ASP 5.37 0 56.454 33.056 46.467 1.00 15.67 MOTA 4886 ASP 710 711 711 35.074 35.217 46.789 45.480 LYS 55.449 1.00 10.56 4887 0 ATOM N 54.802 1.00 16.19 MOTA 4889 CA LYS 0 MOTA 4890 CB LYS 711 54.266 36.661 45.368 1.00 35.75 37.109 37.750 53.743 44.007 1.00 43.76 1.00 49.83 711 4891 LYS n MOTA CG 43.146 4892 4893 54.843 MOTA CD LYS 711 0 711 55.459 38.976 43.819 1.00 54.70 0 LYS CE ATOM 39.514 43.064 1.00 58.93 56.632 711 0 MOTA **4B94** NZ LYS 4898 4899 34.877 C LYS 711 55.753 44.331 1.00 16.03 0 MOTA 55.459 34.001 43.518 1.00 37.96 711 0 LYS00 2.00 1.00 2 MOTA 0 35.560 56.894 44.282 MOTA 4900 N ASP 712 0 ASP 712 57.890 35.330 43.252 4902 CA MOTA 42.927 1.00 75.12 58.655 36.629 712 0 MOTA 4903 CB ASP 1.00 75.41 1.00 77.74 4904 ASP 712 59.362 37.231 44.134 0 CG MOTA 38.013 36.937 712 58.723 44.866 0 OD1 ASP MOTA 4905 60.561 44.341 1.00 84.37 MOTA 4906 OD2 ASP 712 0 1.00 2.00 1.00 77.12 712 58.863 34.189 43.597 4907 ASP MOTA C 60.083 34.366 43.607 0 712 4908 ASP MOTA 0 1.00 27.09 43.866 4909 VAL 713 58.298 33.016 O MOTA N 44.188 713 59.057 31.811 1.00 17.74 VAL MOTA 4911 CA 1.00 2.00 31.579 45.727 4912 59.166 0 MOTA VAL 713 CB 713 713 4913 2.00 CG1 VAL 59.124 30.098 46.069 .1.00 0 MOTA 60.481 32.120 46.224 1.00 2.00 MOTA 4914 CG2 VAL 43.538 43.593 58.319 57.092 30.653 1.00 21.93 0 4915 C VAL 713 MOTA 30.567 29.766 2.00 713 1.00 0 4916 VAL MOTA 0 42.903 59.063 1.00 15.32 714 4917 N LEU MOTA 1.00 10.30 42.249 0 28.642 4919 LEU 714 58.434 MOTA CA 28.147 27.005 1.00 48.90 714 59.285 41.078 0 4920 LEU MOTA CB 40.272 1.00 42.23 4921 4922 4923 714 58.662 LEU MOTA CG 39.909 39.025 1.00 43.72 1.00 42.86 0 27.357 26.753 CD1 LEU 714 57.229 MOTA 59.478 58.224 714 Ω CD2 LEU MOTA 27.530 43.235 1.00 13.23 0 MOTA 4924 C LEU 714 1.00 44.72 1.00 76.36 27.099 27.075 43.445 0 4925 LEU 714 57.096 0 MOTA 715 43.834 0 59.322 MOTA 4926 N GLY 1.00 76.36 25.995 44.800 5**9.26**3 ATOM 4928 CA GLY 715 1.00 76.36 1.00 19.62 0 715 59.630 26.420 46.206 4929 С GLY MOTA 26.974 26.183 46.929 46.583 0 715 58.814 4930 GLY MOTA 1.00 4.87 TRP 716 60.875 MOTA 4931 N 0 716 61.365 26.503 47.918 1.00 4.87 MOTA 4933 CA TRP 48.554 1.00 2.00 0 25.241 61.944 MOTA 4934 CB TRP 716 716 716 48.932 1.00 2.00 TRP 60.884 24.317 4935 CG MOTA 2.00 0 24.529 49.962 1.00 59.936 MOTA 4936 CD2 TRP 23.439 25.541 2.00 0 49.943 1.00 59.050 MOTA 4937 CE2 TRP 716 2.00 0 716 59.750 50.909 1.00 4938 CE3 TRP MOTA 2.00 0 60.562 23.134 48.343 1.00 MOTA 4939 CD1 TRP 716 59.454 57.994 58.715 22.597 48.940 1.00 2.00 0 4940 NE1 TRP 716 MOTA 0 23.334 50.830 1.00 .2.00 716 MOTA 4942 CZ2 TRP 51.783 1.00 2.00 0 25.438 4943 CZ3 TRP 716 MOTA 24.343 27.594 1.00 2.00 0 51.743 716 57.843 4944 TRP MOTA CH2 62.406 63.596 47.954 1.00 4.87 0 4945 TRP 716 ATOM 27.315 47.871 1.00 2.00 4946 TRP 716 MOTA 0 2.00 1.00 0 717 61.968 28.837 48.067 GLY ATOM 4947 Ν 29.937 48.124 1.00 2.00 0 62.911 ATOM 4949 CA GLY 717 49.414 1.00 2.00 30.031 717 63.725 GLY 4950 C MOTA 1.00 2.00 0 63.**44**3 64.752 29.343 50.404 717 4951 GLY MOTA 0 30.880 1.00 4.99 0 49.387 4952 GLU 718 MOTA

> mo:	4054	C.	CI II	710	65.606	31.113	50.538	1.00 8.49	0
MOTA MOTA	4954 4955	CA CB	GLU	718 718	66.980	31.619	50.092	1.00 86.14	ŏ
ATOM	4956	ČĞ	GLU	718	68.026	31.716	51.211	1.00 89.35	0
ATOM	4957	CD	GLU	718	68.627	30.371	51.633	1.00 87.67	0
MOTA	4958	OE1	GLU	718	69.392	30.358	52.624	1.00 95.59 1.00 88.92	0
MOTA	4959	OE2	GLU	718	68.353 64.883	29.336 32.182	50.986 51.340	1.00 88.92	0
MOTA	4960 4961	C O	GLU GLU	718 718	64.597	33.269	50.828	1.00 90.23	Ö
MOTA MOTA	4962	Ŋ	ASN	719	64.560	31.863	52.590	1.00 32.81	ŏ
MOTA	4964	CA	ASN	719	63.848	32.804	53.444	1.00 33.93	0
MOTA	4965	CB	asn	719	63.159	32.083	54.591	1.00 14.30	0
MOTA	4966	CG	ASN	719	62.159	32.965	55.296	1.00 14.30	0
MOTA	4967		ASN	719	62.533 60.879	33.912 32.676	55.983 55.110	1.00 14.30 1.00 14.30	0
MOTA	4968 4971	ND2 C	asn asn	719 719	64.781	33.855	54.007	1.00 33.45	0
MOTA MOTA	4972	Ö	ASN	719	65.825	33.529	54.566	1.00 14.30	ŏ
ATOM	4973	Ň	ASP	720	64.388	35.117	53.878	1.00 42.61	0
ATOM	4975	CA	ASP	720	65.212	36.222	54.351	1.00 47.00	0
MOTA	4976	CB	ASP	720	64.744	37.535	53.706	1.00 77.30	0
MOTA	4977	CG	ASP	720 730	65.025 66.104	37.574 38.071	52.200 51.801	1.00 92.85 1.00 92.39	0
MOTA	4978 4979	OD1 OD2	ASP	720 720	64.172	37.094	51.416	1.00 90.51	ő
MOTA MOTA	4980	C	ASP	720	65.328	36.352	55.869	1.00 39.89	ŏ
MOTA	4981	ŏ	ASP	720	66.255	36.986	56.370	1.00 74.64	Ō
MOTA	4982	N	ARG	721	64.411	35.728	56.602	1.00 13.83	0
MOTA	4984	CA	ARG	721	64.446	35.759	58.060	1.00 12.18	0
MOTA	4985	CB	ARG	721	63.262 61.946	35.007	58.649 58.572	1.00 15.37 1.00 7.28	0
MOTA	4986	CC	ARG ARG	721 721	60.950	35.693 34.836	59.300	1.00 7.28	ő
MOTA MOTA	4987 4988	CD NE	ARG	721 721	59.593	35.306	59.103	1.00 8.88	ő
MOTA	4990	CZ	ARG	721	59.051	36.315	59.766	1.00 9.83	Ö
ATOM	4991		ARG	721	59.751	36.965	60.690	1.00 8.95	0
MOTA	4994	NH2	ARG	721	57.809	36.682	59.485	1.00 8.62	0
MOTA	4997	C	ARG	721	65.710	35.093	58.592	1.00 8.44	0
MOTA	4998	0	ARG	721 722	65. 98 2 66. 44 9	35.142 34.437	59.798 57.697	1.00 10.25 1.00 4.85	ő
MOTA MOTA	4999 5001	N CA	GLY GLY	722	67.668	33.752	58.083	1.00 4.85	ŏ
ATOM	5002	Ç.	GLY	722	67.392	32.344	58.592	1.00 4.85	0
MOTA	5003	ŏ	GLY	722	68.305	31.655	59.052	1.00 76.68	0
ATOM	5004	N	VAL	72 3	66.134.	31.918	58.508	1.00 10.85	0
MOTA	5006	CA	VAL	723	65.717	30.591	58.958	1.00 10.85 1.00 2.00	0
MOTA	5007	CB	VAL	72 3 72 3	65. 25 9 66. 44 9	30. 574 30. 40 0	60.471 61.386	1.00 2.00 1.00 2.00	ŏ
MOTA MOTA	5008 5009	CG1 CG2	VAL	723 723	64.504	31.858	60.825	1.00 2.00	ŏ
ATOM	5010	C	VAL	72 3	64.546	30.100	58.105	1.00 10.85	0
ATOM	5011	ō	VAL	72 3	63 .66 7	30.883	57.714	1.00 2.00	0
MOTA	5012	N	SER	724	64.541	28.802	57.824	1.00 42.15	0
MOTA	5014	CA	SER	724	63.479	28.193	57.040	1.00 42.15 1.00 2.00	0
MOTA	5015	CB	SER	724	62.127	28.609 27.808	57.623 57.106	1.00 2.00 1.00 2.00	0
MOTA ATOM	5016 5018	og C	Ser Ser	724 724	61.077	28.588	55.561	1.00 42.15	č
ATOM	5019	Ö	SER	724	64.680	28.820	55.049	1.00 2.00	0
ATOM	5020	N	PHE	725	62.454	28.643	54.863	1.00 2.00	0
MOTA	5022	CA	PHE	725	62.464	29.007	53.457	1.00 2.00	0
MOTA	5023	CB	PHE	725	62.461	27.779	52.541	1.00 2.00	0
ATOM	5024	CG	PHE	725 725	62.891 62.0 4 7	26.551 25.916	53.205 54.089	1.00 2.00 1.00 2.00	0
MOTA MOTA	5025 5026	CD1		725 725	64.151	26.044	52.984	1.00 2.00	ŏ
ATOM	5027	CE1	PHE	725	62.458	24.791	54.750	1.00 2.00	Ō
MOTA	5028	CE2	PHE	725	64.578	24.906	53.646	1.00 2.00	0
MOTA	5029	CZ	PHE	725	63.733	24.280	54.530	1.00 2.00	0
ATOM	5030	C	PHE	725	61.222	29.787	53.146	1.00 2.00	0
ATOM	5031	0	PHE	725	60.382	30.040	54.009	1.00 2.00	0
ATOM	5032	N	THR	726	61.132	30.162 30.877	51.886 51.357	1.00 34.54 1.00 30.82	0
ATOM	5034	CA	THR THR	726 726	60. 009 60. 46 8	32.154	50.661	1.00 30.82	Ö
ATOM ATOM	5035 5036	CB OG1	THR	726	61.880	32.098	50.406	1.00 2.00	Ö
ATOM	5038		THR	726	60.222	33.327	51.561	1.00 2.00	Ō

MOTA	5039	С	THR	726	59.450	29.866	50.382	1.00 34.20	0
ATOM	5040	0	THR	726	60.201	29.208	49.684	1.00 2.00	0
MOTA	5041	N	PHE	727	58.144	29.685	50.375	1.00 2.00	0
	5043	CA	PHE	727	57.555	28.712	49.483	1.00 2.00	ő
ATOM				727	56.852	27.612	50.275	1.00 9.97	
MOTA	5044	CB	PHE				51.105	1.00 15.33	0
ATOM	5045	CG	PHE	727	55.698	28.096			0
MOTA	5046	CD1	PHE	727	54.394	27.922	50.666	1.00 9.89	0
ATOM	5047	CD2	PHE	727	55.916	28.726	52.323	1.00 15.65	0
ATOM	5048	CE1	PHE	727	53.338	2B.360	51.418	1.00 11.89	0
ATOM	5049	CE2	PHE	727	54.852	29.171	53.087	1.00 6.87	0
MOTA	5050	cz	PHE	727	53.563	28.986	52.631	1.00 11.72	Ö
		c	PHE	727	56.580	29.398	48.553	1.00 2.00	ŏ
MOTA	5051				55.848	30.312	48.982	1.00 18.77	ő
MOTA	5052	0	PHE	727					
MOTA	5053	N	GLY	728	56. 5 76	28.956	47.289	1.00 13.14	0
MOTA	5055	CA	GLY	728	55.709	29.542	46.277	1.00 12.12	0
MOTA	5056	С	GLY	728	54.348	28.894	46.174	1.00 13.69	0
ATOM	5057	0	GLY	728	54.062	27.913	46.862	1.00 2.00	0
ATOM	5058	N	ALA	729	53.513	29.436	45.292	1.00 33.39	0
	5060	CA	ALA	729	52.161	28.916	45.080	1.00 33.11	0
MOTA				729	51.375	29.862	44.200	1.00 16.31	ŏ
MOTA	5061	CB	ALA			27.507	44.485	1.00 31.34	ŏ
MOTA	5062	C	ALA	729	52.139				
ATOM	5063	0	ALA	72 9	51.143	26.796	44.600	1.00 16.31	0
ATOM	5064	N	GLU	73 0	53.221	27.105	43.831	1.00 22.53	0
MOTA	5066	CA	GLU	730	53.284	25.761	43.281	1.00 26.87	0
MOTA	5067	CB	GLU	730	54.622	25.551	42.570	1.00 59.75	0
ATOM	5068	CG	GLU	730	54.893	24.117	42.142	1.00 62.63	0
		CD	GLU	730	56.138	23.984	41.283	1.00 68.00	0
MOTA	5069			730	57.213	23.624	41.822	1.00 74.97	ō
MOTA	5070	OE1	GLU			24.238	40.063	1.00 67.31	ŏ
ATOM	5071	OE2	GLU	730	56.034				ő
MOTA	5072	C	GLU	730	53.140	24.781	44.446	1.00 24.85	
ATOM	5073	0	GLU	730	52.285	23.899	44.425	1.00 58.27	0
ATOM	5074	N	VAL	731	53. 95 8	24.989	45.477	1.00 24.13	0
ATOM	5076	CA	VAL	731	53.985	24.157	46.679	1.00 19.42	0
ATOM	5077	CB	VAL	731	55.079	24.628	47.645	1.00 19.11	0
ATOM	5078	CG1	VAL	731	55.159	23.699	48.824	1.00 19.11	0
			VAL	731	56.412	24.691	46.942	1.00 19.11	0
MOTA	5079	CG2	A VIT	1.01		2			
MOTA		_	7777	721		24 165	47 423	1 00 16.89	0
	5080	C	VAL	731	52.659	24.165	47.423	1.00 16.89	0
ATOM	5081	0	VAL	731	52.659 52.210	23.128	47.900	1.00 19.11	0
ATOM ATOM	5081 5082	N O	VAL VAL	731 732	52.659 52.210 52.035	23.128 25.332	47.900 47.525	1.00 19.11 1.00 15.54	0
	5081	0	VAL	731 732 732	52.659 52.210 52.035 50.750	23.128 25.332 25.449	47.900 47.525 48.209	1.00 19.11 1.00 15.54 1.00 15.54	000
ATOM	5081 5082	N O	VAL VAL	731 732 732 732	52.659 52.210 52.035 50.750 50.254	23.128 25.332 25.449 26.898	47.900 47.525 48.209 48.240	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17	0 0 0
ATOM ATOM ATOM	5081 5082 5084	O N CA	VAL VAL VAL	731 732 732	52.659 52.210 52.035 50.750	23.128 25.332 25.449 26.898 26.985	47.900 47.525 48.209 48.240 48.962	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17 1.00 20.17	0 0 0 0
MOTA MOTA MOTA	5081 5082 5084 5085 5086	O N CA CB CG1	VAL VAL VAL VAL	731 732 732 732	52.659 52.210 52.035 50.750 50.254	23.128 25.332 25.449 26.898	47.900 47.525 48.209 48.240	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17	0 0 0 0
ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087	O N CA CB CG1 CG2	VAL VAL VAL VAL VAL VAL	731 732 732 732 732 732 732	52.659 52.210 52.035 50.750 50.254 48.907	23.128 25.332 25.449 26.898 26.985	47.900 47.525 48.209 48.240 48.962	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17 1.00 20.17	0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088	O N CA CB CG1 CG2 C	VAL VAL VAL VAL VAL VAL	731 732 732 732 732 732 732 732	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653	23.128 25.332 25.449 26.898 26.985 27.764 24.609	47.900 47.525 48.209 48.240 48.962 48.902 47.554	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17	0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089	O N CA CB CG1 CG2 C	VAL VAL VAL VAL VAL VAL VAL	731 732 732 732 732 732 732 732 732	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791	47.900 47.525 48.209 48.240 48.962 48.902 47.554 48.222	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17	000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5090	O N CA CB CG1 CG2 C	VAL VAL VAL VAL VAL VAL VAL	731 732 732 732 732 732 732 732 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822	47.900 47.525 48.209 48.240 48.962 48.902 47.554 48.222 46.253	1.00 19.11 1.00 15.54 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5090 5092	O N CA CB CG1 CG2 C O N CA	VAL VAL VAL VAL VAL VAL VAL ALA	731 732 732 732 732 732 732 732 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.611 49.437 48.408	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29	000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5090 5092 5093	O N CA CB CG1 CG2 C O N CA CB	VAL VAL VAL VAL VAL VAL VAL ALA ALA	731 732 732 732 732 732 732 732 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.653 49.011 49.437 48.408 48.260	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33	0000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5090 5092 5093	O N CA CB CG1 CG2 C O N CA	VAL VAL VAL VAL VAL VAL VAL ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618	47.900 47.525 48.209 48.240 48.962 48.902 47.554 48.222 46.253 45.501 44.131 45.390	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 16.29 1.00 16.29	00000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5090 5092 5093	O N CA CB CG1 CG2 C O N CA CB	VAL VAL VAL VAL VAL VAL VAL ALA ALA	731 732 732 732 732 732 732 732 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795	47.900 47.525 48.209 48.240 48.962 48.902 47.554 48.222 46.253 45.501 44.131 45.390 45.349	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78	0000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5099 5099 5099 5095	O N CA CB CG1 CG2 C O N CA CB C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.348	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00	00000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5087 5088 5089 5099 5099 5099 5099 5099	O N CA CB CG1 CG2 C O N CA CB C O N CA CB C O N	VAL VAL VAL VAL VAL VAL VAL VAL ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795	47.900 47.525 48.209 48.240 48.962 48.902 47.554 48.222 46.253 45.501 44.131 45.390 45.349	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.33 1.00 16.29 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5087 5088 5089 5090 5092 5093 5094 5098	O N CA CG2 C O N CA CB C O N CA CB C C	VAL VAL VAL VAL VAL VAL VAL ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.348	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03	0000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5088 5089 5099 5099 5099 5099 5099	O N CA CG2 C O N CA CB C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.012	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287 20.905 20.936	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.348 45.259 45.015	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03	000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5088 5089 5099 5099 5099 5099 5099 5100	O N CA CB CG1 CG2 C O N CA CB C C O N CA CB C C C C C C C C C C C C C C C C C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.012 52.507	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287 20.905 20.936 20.244	47.900 47.525 48.209 48.962 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.349 45.349 45.359 45.259 43.759	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55	0000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5088 5089 5099 5099 5099 5099 5099 5101	O N CA CB CG O N CA CB C CD CCD	VAL VAL VAL VAL VAL ALA ALA ALA ALA LYS LYS LYS	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.012 52.507 52.696	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287 20.905 20.936 20.244 21.212	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.259 45.259 45.259	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13	000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5088 5089 5099 5099 5099 5099 5101 5102	O N CA CB CG O N CA CB C CD CC CC	VAL VAL VAL VAL VAL VAL ALA ALA ALA LYS LYS LYS LYS	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.611 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.010 52.507 52.696 53.613	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.109 24.717 22.618 21.795 20.905 20.936 20.936 20.244 21.212 20.608	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.259 45.015 42.594 41.521	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 25.55 1.00 35.13 1.00 40.96	00000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5088 5088 5088 5088 5088 5099 5099 5099	O N CA CB CG2 C O N CA CCC CC CC CC CC NZ	VAL VAL VAL VAL VAL VAL ALA ALA ALA LYS LYS LYS LYS LYS	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.012 52.507 53.613 55.006	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.109 24.717 22.618 21.795 22.287 20.905 20.936 20.244 21.212 20.608 20.311	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.349 45.259 45.015 43.759 441.521 42.001	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 46.09	00000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 5085 5086 5088 5089 5099 5099 5099 5099 5101 5102	O N CA CB CG O N CA CB C CD CC CC	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.012 52.507 52.696 53.613 55.006 50.222	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 20.905 20.936 20.244 21.212 20.608 20.311 20.146	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.348 45.259 45.015 43.759 42.594 41.521 46.578	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 46.09 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5088 5088 5088 5088 5088 5099 5099 5099	O N CA CB CG2 C O N CA CCC CC CC CC CC NZ	VAL VAL VAL VAL VAL VAL ALA ALA ALA LYS LYS LYS LYS LYS	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.260 48.776 49.996 50.499 52.012 52.696 53.616 50.222 49.995	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.295 20.905 20.936 20.244 21.212 20.608 20.146 18.926	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.348 45.259 45.015 43.759 42.599 41.521 42.594 41.521 42.578 46.591	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 15.54 1.00 16.29 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 40.96 1.00 2.00 1.00 16.68	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50886 50887 50889 50990 50992 50998 50998 50998 51002 51002 51007 5108	O N CAB CG12 C O N CAB C C C C C C C C C C C C C C C C C C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.012 52.507 52.696 53.613 55.006 50.222	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 20.905 20.936 20.244 21.212 20.608 20.311 20.146	47.900 47.525 48.209 48.962 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.349 45.259 42.599 42.594 41.521 42.001 46.591 47.670	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 40.96 1.00 2.00 1.00 2.00 1.00 36.96	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50886 50887 5089 50990 50992 50998 50998 50998 5100 5100 5100 5100 5100 5100 5100 510	O N CAB CGG C C O N CAB CG C C C O N CAB CG C C C O N CAB C C C C C C C C C C C C C C C C C C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.260 48.776 49.996 50.499 52.012 52.696 53.616 50.222 49.995	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.295 20.905 20.936 20.244 21.212 20.608 20.146 18.926	47.900 47.525 48.209 48.962 48.962 47.554 48.222 46.253 45.501 44.131 45.349 45.349 45.349 45.015 43.759 42.594 41.521 42.001 46.578 46.578 47.670 49.047	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 15.54 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 46.09 1.00 2.00 1.00 2.00 1.00 33.73	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50885 50889 5089 50992 50993 50993 50996 5100 5100 5100 5100 5100 5100 5100 510	O N CAB CGC C O N CAB CGC C C O N CAB CGC C O N CAB C C C C C C C C C C C C C C C C C C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 733 733 733 733 734 734 734 734 734 734	52.659 52.210 52.035 50.750 50.254 48.207 49.653 49.011 49.437 48.260 48.703 47.776 49.996 52.012 52.507 52.696 53.613 55.002 49.995 50.225 50.2263 50.036	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287 20.905 20.244 21.212 20.608 20.311 20.146 18.926 20.915	47.900 47.525 48.209 48.962 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.349 45.259 42.599 42.594 41.521 42.001 46.591 47.670	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 40.96 1.00 2.00 1.00 2.00 1.00 36.96	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5084 50885 50887 50889 50992 50993 50995 51003 51003 51003 51103 51103 51103 51112	O N CAB CGC CO N CAB CCC CO N CAB CCC CO N CAB CCC CO N CAB CCC CO N CAB	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 733 733 733 733 734 734 734 734 734 734	52.659 52.210 52.035 50.750 50.254 48.907 51.653 49.011 49.437 48.408 48.703 47.776 49.999 52.012 52.696 53.613 55.006 50.222 49.995 50.036 50.036 50.036	23.128 25.332 25.449 26.8985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287 20.905 20.244 21.212 20.608 20.311 20.146 18.926 20.915 20.478 21.554	47.900 47.525 48.209 48.962 48.962 47.554 48.222 46.253 45.501 44.131 45.349 45.349 45.349 45.015 43.759 42.594 41.521 42.001 46.578 46.578 47.670 49.047	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 15.54 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 46.09 1.00 2.00 1.00 2.00 1.00 33.73	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50885 50887 50889 50992 50993 50996 50999 51003 51003 51003 51003 51103 51112 51113	ONCABCCONCABCCCCCCCCCCCCCCCCCCCCCCCCCCCC	VAL VAL VAL VAL VAL ALA ALA ALA ALA LYS LYS LYS LYS LYS PHE PHE PHE	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.400 48.703 47.776 49.996 50.499 52.507 53.613 50.222 49.995 50.2696 50.202 50.606 50.320	23.128 25.332 25.449 26.8985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 22.287 20.905 20.936 20.244 21.212 20.608 20.311 20.146 18.926 20.915 20.915 21.554 21.323	47.900 47.525 48.240 48.962 48.952 47.554 48.253 45.501 44.131 45.349 45.349 45.349 45.259 441.521 42.594 41.521 42.591 42.670 46.591 47.670 49.991 51.465	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 16.29 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 46.09 1.00 16.68 1.00 36.96 1.00 33.73 1.00 8.60 1.00 8.60	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 5088 5088 5088 5088 5089 5099 5099 5099	ON CAB CGC CON CAB CCC CON CAB CCC CON CAB CCC CON CAB CCC CCC CON CON CCC CCC CCC CCC CCC CCC	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 732 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.776 49.996 50.499 52.507 52.696 50.222 49.995 50.036 50.0320 50.955	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 20.905 20.936 20.244 21.212 20.311 20.146 18.926 20.478 21.554 21.554 21.523 20.298	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.349 45.348 45.259 45.015 43.759 42.591 42.591 46.578 46.591 47.670 49.991 51.465 52.169	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 20.17 1.00 15.54 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 2.00 1.00 25.55 1.00 35.13 1.00 46.09 1.00 46.09 1.00 46.09 1.00 16.68 1.00 36.96 1.00 36.96 1.00 8.60 1.00 8.60	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50886 50889 50990 50990 50990 50990 5100 5110 5110	ON CAB CGC ON CAB CGC CC ON CAB CGC CC CON CAB CGC CC CON CON CON CCC CCC CCC CCC CCC CCC	VAL	731 732 732 732 732 732 733 733 733 733 734 734 734 734 734 734	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.408 48.260 48.703 47.776 49.996 50.499 52.612 52.507 52.696 50.222 49.995 50.263 50.606 50.205 50.606 50.955 49.412	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.2795 20.905 20.936 20.244 21.212 20.608 20.146 18.926 20.478 21.554 21.554 21.523 20.298 22.135	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.348 45.259 45.015 43.759 42.599 41.521 42.591 41.521 42.591 41.521 42.591 41.591 41.591 41.591 41.591 41.670 49.991 51.465 52.169 52.142	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 16.29 1.00 16.29 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 46.09 1.00 46.09 1.00 16.68 1.00 36.96 1.00 33.73 1.00 8.60 1.00 8.60 1.00 8.60	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50886 50889 50990 50990 50998 50998 5100 5110 5110 51112 51114 51116	ON CAB CG12 CON CAB CCC CON CAB CCC CCC CON CAB CCC CON CAB CCC CCC CCC CCC CCC CCC CCC CCC CCC	VAL	731 732 732 732 732 732 733 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.260 48.260 48.260 50.499 52.012 52.507 52.696 53.610 50.222 49.995 50.263 50.363 50.365 50.325 49.412 50.690	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.295 20.905 20.936 20.244 21.212 20.608 20.311 20.478 21.554 21.554 21.323 22.135 20.086	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.348 45.259 45.015 43.759 42.591 42.001 46.591 47.670 49.991 51.469 52.142 53.512	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 15.54 1.00 16.29 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 2.00 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 40.96 1.00 16.68 1.00 36.96 1.00 33.73 1.00 8.60 1.00 8.60 1.00 8.60 1.00 8.60	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50885 50889 50992 50993 50998 50998 5100 51102 51103 51112 51113 51114 51117	O N CAB CG12 C O N CAB C C C C C C C C C C C C C C C C C C	VAL VAL VAL VAL VAL VAL ALA ALA ALA ALA	731 732 732 732 732 732 733 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.437 48.408 48.260 48.776 49.996 50.499 52.696 53.610 50.222 49.995 50.2696 50.325 50.606 50.325 49.413	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.795 20.905 20.936 20.244 21.212 20.608 20.311 20.478 21.554 21.554 21.323 20.298 21.323 20.298 21.929	47.900 47.525 48.209 48.962 48.962 47.554 48.222 46.253 45.301 44.131 45.390 45.349 45.349 45.349 45.259 42.591 42.591 42.501 46.591 47.670 49.991 51.465 52.165 53.512 53.483	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 20.17 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 17.78 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 40.96 1.00 40.96 1.00 40.96 1.00 33.73 1.00 40.96 1.00 36.96 1.00 8.60 1.00 8.60 1.00 8.60 1.00 8.60 1.00 8.60	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5081 5082 50884 50886 50889 50990 50990 50998 50998 5100 5110 5110 51112 51114 51116	ON CAB CG12 CON CAB CCC CON CAB CCC CCC CON CAB CCC CON CAB CCC CCC CCC CCC CCC CCC CCC CCC CCC	VAL	731 732 732 732 732 732 733 733 733 733 733	52.659 52.210 52.035 50.750 50.254 48.907 51.272 49.653 49.011 49.437 48.260 48.260 48.260 50.499 52.012 52.507 52.696 53.610 50.222 49.995 50.263 50.363 50.365 50.325 49.412 50.690	23.128 25.332 25.449 26.898 26.985 27.764 24.609 23.791 24.822 24.109 24.717 22.618 21.295 20.905 20.936 20.244 21.212 20.608 20.311 20.478 21.554 21.554 21.323 22.135 20.086	47.900 47.525 48.209 48.240 48.962 47.554 48.222 46.253 45.501 44.131 45.390 45.348 45.259 45.015 43.759 42.591 42.001 46.591 47.670 49.991 51.469 52.142 53.512	1.00 19.11 1.00 15.54 1.00 20.17 1.00 20.17 1.00 15.54 1.00 15.54 1.00 16.29 1.00 16.29 1.00 16.29 1.00 17.33 1.00 16.29 1.00 2.00 1.00 2.00 1.00 2.00 1.00 23.03 1.00 25.55 1.00 35.13 1.00 40.96 1.00 40.96 1.00 16.68 1.00 36.96 1.00 33.73 1.00 8.60 1.00 8.60 1.00 8.60 1.00 8.60	000000000000000000000000000000000000000

ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	55555555555555555555555555555555555555	CCONCECCNCONCECCNCONCECCNCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCECCOOCCONCE	LEU LEU ASP ASP ASP ASP	77777777777777777777777777777777777777	48.5451 48.731 46.739 45.7314 46.2997 45.314 45.747 46.638 47.7603 46.638 47.408 46.638 47.408 46.638 47.408 46.638 47.408 46.638 47.408 46.638 47.418 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 46.638 47.423 47.423 48.609 48.609 48.609 48.610 42.620 41.900 42.620 42.620 42.620 43.762 41.900 42.620 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 43.774 44.778 44.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46.620 46	20.256 19.181 21.288 22.451 23.760 24.883 29.906 19.215 19.906 18.037 17.162 19.215 17.162 16.046 17.162 16.046 17.202 16.046 17.202 16.316	49.301 49.079 49.48.689 50.2970 49.48.689 50.2970 49.48.689 50.298 49.48.689 50.2888 49.1037 40.45.268 40.2888	1.00 34.15 1.00 8.60 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 25.73 1.00 69.97 1.00 68.65 1.00 76.42 1.00 76.45 1.00 76.42 1.00 76.42 1.00 10.83 1.00 10.	
ATOM ATOM ATOM ATOM	5198 5199 5200 5201	CB CG							



		CB GLU	752	48.430	46.435	54.190	1.00 39.24	0
MOTA	5287 5288	CB GLU	752	49.867	46.349	53.628	1.00 75.32	0
MOTA MOTA	5289	CD GLU	752	50.962	46.804	54.611	1.00 75.31	0
ATOM	5290	OE1 GLU	752	52.007	46.121	54.687	1.00 74.90 1.00 91.76	0
ATOM	5291	OE2 GLU	752	50.792	47.837	55.300 55.913	1.00 2.00	Ö
ATOM	5292	c Gro	752	46.744 46.326	45.716 46.469	56.796	1.00 38.94	Ö
MOTA	5293	O GLU	752 753	45.968	44.860	55.259	1.00 31.76	ŏ
ATOM	5294	N ASP	753 753	44,527	44.776	55.491	1.00 34.73	Ŏ
MOTA	5296 5297	CA ASP	753	43.787	44.470	54.170	1.00 79.26	0
MOTA MOTA	5298	CG ASP	753	44.539	43.487	53.251	1.00 85.10	0
ATOM	5299	OD1 ASP	753	44.040	43.240	52.130	1.00 88.01 1.00 92.40	0
ATOM	5300	OD2 ASP	753	45.613	42.960 43.853	53.619 56.606	1.00 92.40 1.00 31.43	0
MOTA	5301	C ASP	753 753	44.028 42.824	43.759	56.835	1.00 64.64	ő
MOTA	5302	O ASP N GLY	754	44.940	43.188	57.309	1.00 8.60	Õ
MOTA	5303 5305	N GLY		44,535	42.309	58.393	1.00 5.10	0
MOTA MOTA	5306	C GLY	754	44.577	40.849	58.007	1.00 3.66	0
ATOM	5307	O GLY	754	44.781	39.959	58.853	1.00 2.00	0
MOTA	5308	n TY R		44.361	40.611	56.718	1.00 27.09 1.00 27.09	0
ATOM	5310	CA TYR		44.388	39.282 38.632	56.129 56.128	1.00 27.03	Ö
MOTA	5311	CB TYR		43.004 41.976	39.338	55.279	1.00 25.47	ŏ
MOTA	5312 5313	CG TYR		41.236	40.393	55.797	1.00 23.65	0
MOTA MOTA	5314	CE1 TYR		40.276	41.049	55.034	1.00 28.13	0
ATOM	5315	CD2 TYR		41.736	38.949	53.964	1.00 30.24	0
ATOM	5316	CE2 TYR		40.774	39.601	53.187	1.00 27.11 1.00 32.27	0
MOTA	5317	CZ TYR		40.046	40.653 41.311	53.732 52.996	1.00 28.90	Ö
MOTA	5318	OH TYR		39.086 44.844	39.530	54.713	1.00 27.09	ō
MOTA	5320 5321	C TYR		44.772	40.654	54.238	1.00 25.53	0
MOTA MOTA	5322	N GLU		45.288	38.483	54.035	1.00 8.71	0
MOTA	5324	CA GLU		45.787	38.617	52.676	1.00 8.71 1.00 5.76	0
MOTA	5325	CB GLU		47.170	39.251	52.740 51.436	1.00 5.76 1.00 16.72	ő
MOTA	5326	CG GLU		47.832 49.127	39.557 40.303	51.430	1.00 23.34	Ŏ
MOTA	5327	OE1 GLU		49.074	41.554	51.801	1.00 25.72	0
MOTA MOTA	5328 5329	OE1 GLU		50.194	39.636	51.751	1.00 25.15	0
ATOM	5330	C GLU		45.853	37.240	52.044	1.00 8.71	0
ATOM	5331	O GLU		46.400	36.306	52.638	1.00 6.14 1.00 28.33	0
ATOM	5332	N PHE		45.269	37.105 35.822	50.858 50.174	1.00 28.33	ŏ
ATOM	5334	CA PHE		45.270 44.055	35.683	49.253	1.00 2.00	0
MOTA MOTA	5335 5336	CB PHE		42.748	35.612	49.982	1.00 2.00	0
ATOM	5337	CD1 PHE		41.961	36.747	50.137	1.00 2.00	0
ATOM	5338	CD2 PHE		42.306	34.407	50.535	1.00 2.00 1.00 2.00	0
MOTA	5339	CE1 PHE		40.751	36.695 34.337	50.833 51.238	1.00 2.00 1.00 2.00	ŏ
MOTA	5340	CE2 PHE		41.092 40.315	35.489	51.385	1.00 2.00	Ö
MOTA	5341	CZ PHE		46.547	35.639	49.387	1.00 28.33	0
MOTA MOTA	5342 5343	O PHE		47.220	36.602	49.033	1.00 2.00	0
ATOM	5344	N PHE		46.893	34.387	49.150	1.00 2.00	0
ATOM	5346	CA PHE	758	48.075	34.050	48.397	1.00 2.00 1.00 10.52	ŏ
MOTA	5347	CB PHE		49.241	33.789 33.263	49.317 48.618	1.00 7.54	ō
MOTA	5348	CG PHI		50.450 51.404	34.131	48.100	1.00 7.91	0
MOTA MOTA	5349 5350	CD1 PHE		50.654	31.895	48.492	1.00 7.80	0
ATOM	5351	CE1 PH		52.546	33.642	47.467	1.00 11.14	0
ATOM	5352	CE2 PH	758	51.796	31.398	47.858	1.00 11.57	0
ATOM	535 3	CZ PHI		52.741	32.272	47.347 47.636	1.00 10.83 1.00 2.00	0
MOTA	5354	C PH		47.725 46.826	32.786 32.034	48.042	1.00 2.00	Ö
MOTA	5355	O PHE		48.415	32.573	46.518	1.00 17.88	0
ATOM ATOM	5356 5358	N ALA		48.196	31.407	45.673	1.00 17.63	0
ATOM	5359	CB ALA	759	48.767	30.155	46.334	1.00 2.00	0
ATOM	5360	C ALA	759	46.733	31.180	45.314	1.00 19.54 1.00 2.00	0
ATOM	5361	O ALA		46.187	30.106	45.547 44 780	1.00 2.00	٥
ATOM	5362	H LY	760	46.086	32.205	44 /00	1.55 9.90	3

ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5365 53667 53667 53667 5377 5377 53377 53377 53377 53381 53381 53391 533	CCBCCCNC ON CCBCCCNNC ON CCBCCCONC ON CONCONCONCONCONCONCONCONCONCONCONCONCONC	LYS LYS LYS LYS LYS LYS LYS ARG ARG ARG ARG ARG ARG ARG ARG GLN GLN GLN GLN GLN GLN GLN LEU LEU	760 760 760 760 760 761 761 761 761 762 762 762 762 763 763	44.699 44.699 45.654 45.654 45.6843 44.506 44.621 43.715 42.953 43.751 42.869 41.399 41.399 41.399 38.798 39.988 43.145 42.502 44.359 45.167 44.502 45.309 45.111 46.236 45.278 45.631 46.354	32.076 31.068 31.316 30.068 29.540 28.293 31.659 30.702 32.379 32.154 33.375 34.493 35.785 36.642 37.963 30.844 30.639 27.794 26.959 25.745 27.596 27.893 30.053	44.341 43.184 42.065 40.699 39.896 45.456 46.5721 47.256 46.5721 47.274 46.458 47.334 46.6898 47.767 46.898 47.767 48.559 47.553 46.196 45.097 44.476 49.585 47.46.196 49.585 47.775 49.585 49.585 47.559 47.559 47.46.196 49.585	1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00	64.10 64.10 64.10 64.10 64.10 64.10 64.10 64.10 64.10 64.10 64.10 8.86 38.86 38.86 38.86 38.86 38.86 38.86 16.83 38.86 16.83 38.86 16.83 38.86 16.83 38.86 3	000000000000000000000000000000000000000
MOTA MOTA	5407 5408	CB CG	Leu Leu	763 763	47.882 48.8 4 7	30.319 30.427	51.127 52.335		13.68 13.68	0
ATOM	5409	CD1	LEU	763	50.117	29.641	52.066	1.00	13.68	0
MOTA	5410 5411	CD2 C	LEU	763 763	49.200 45.893	31.871 31.705	52.633 51.809	1.00 1.00	13.68 2.00	0
ATOM	5412	Ö	LEU	763	45.654	32.595	50.981	1.00	13.68	0
MOTA	5413	N	VAL	764	45.741	31.869	53.118		13.21	0
MOTA	5415	CA	VAL	764 764	45.340 43.825	33.140 33.165	53.680 53.953	1.00	13.65 2.00	0
MOTA MOTA	5416 5417	CB CG1	VAL VAL	764	43.452	32.097	54.933	1.00	2.00	ŏ
MOTA	5418	CG2	VAL	764	43.404	34.521	54.434	1.00	2.00	0
ATOM ATOM	5419 5420	C O	VAL VAL	764 764	46.159 46.396	33.363 32.437	54.955 55.737	1.00	18.12	0
ATOM	5421	N	THR	765	46.646	34.587	55.111	1.00	2.00	0
MOTA	5423	CA	THR	765	47.453	34.992	56.254 55.798	1.00 1.00	2.00	0
MOTA MOTA	5424 5425	CB OG1	THR THR	7 65 765	48.7 3 1 49.474	35.660 34.740	54.991	1.00	2.00	ő
ATOM	5427	CG2	THR	765	49.542	36.119	56.973	1.00	2.00	0
ATOM	5428	C	THR	765	46.675 46.201	36.024 37.016	57.030 56. 4 54	1.00	2.00 2.00	0
ATOM ATOM	5429 5430	И О	THR L E U	765 766	46.549	35.797	58.333	1.00	5.65	ŏ
MOTA	5432	CA	LEU	766	45.832	36.717	59.205	1.00	5.65	0
ATOM	5433	CB	LEU	766 766	44.848 43.964	35.943 34.922	60.059 59.372	1.00	2.00 2.00	0
MOTA MOTA	5434 5435	CG CD1	LEU LEU	766	43.703	33.794	60.327	1.00	2.00	0
MOTA	5436		LEU	766	42.672	35.581	58.943	1.00	2.00 5.65	0
MOTA MOTA	5437 5438	CO	LEU LEU	766 766	46.826 47.864	37. 38 2 36. 79 0	60.142 60.47B	1.00	2.00	Ö
MOTA	5439	N	PHE	767	46.520	38.605	60.554	1.00	2.00	0
MOTA	5441	CA	PHE	767	47.342	39.320	61.530 60.864	1.00	2.00	0
MOTA MOTA	5442 5443	CB CG	PHE PHE	767 767	48.259 49.494	40.328 40.612	61.649	1.00	2.00	ő
MOTA	5444	CD1	PHE	767	50.523	39.678	61.695	1.00	2.00	0
ATOM	5445	CD2	PHE	767	49.647 51.689	41.811 39.935	62.317 62.389	1.00 1.00	2.00	0
ATOM ATOM	5446 5447		PHE PHE	767 767	50.813	42.078	63.018	1.00	2.00	ő
MOTA	5448	CZ	PHE	767	51.838	41.134	63.050	1.00	2.00	0
ATOM	5449	С	PHE	7 67	46.325	40.033	62.418	1.00	2.00	0

AROM 5450 O PHE 767 AROM 5451 N SER 768 ASSET			_			45 0E7	41.184	62.171	1.00 2.00	0
ANDION 5453 CA SER 768										
ATOM \$455 CS SER 768 \$43,003 37,635 64,175 1,000 2,000 0										
ATOM 5455 OC SER 768 43.903 37.635 64.175 1.00 2.00 0 ATOM 5458 O SER 768 45.261 40.875 65.336 1.00 2.00 0 ATOM 5458 N ALA 769 44.639 41.911 65.474 1.00 30.20 0 ATOM 5458 N ALA 769 44.639 41.911 65.474 1.00 30.20 0 ATOM 5461 CA ALA 769 44.639 41.911 65.474 1.00 30.20 0 ATOM 5462 CB ALA 769 44.639 41.911 65.474 1.00 30.20 0 ATOM 5463 C ALA 769 45.410 42.2454 66.428 1.00 27.69 0 ATOM 5463 C ALA 769 44.639 41.92 66.428 1.00 27.69 0 ATOM 5466 N PRO 770 45.410 20.00 0 ATOM 5466 CD PRO 770 47.556 42.2904 56.989 1.00 11.84 0 ATOM 5466 CD PRO 770 47.556 42.2904 56.989 1.00 1.00 0 ATOM 5468 CD PRO 770 47.556 42.904 56.939 1.00 1.00 0 ATOM 5468 CD PRO 770 48.913 43.329 55.392 1.00 1.00 0 ATOM 5468 CD PRO 770 48.913 43.329 55.392 1.00 1.00 0 ATOM 5467 N PRO 770 47.227 46.584 64.825 1.00 14.07 0 ATOM 5467 CD PRO 770 48.913 43.329 55.392 1.00 1.00 0 ATOM 5467 CD PRO 770 48.913 43.329 55.392 1.00 1.00 0 ATOM 5467 CD PRO 770 48.913 43.329 65.392 1.00 1.00 0 ATOM 5467 CD PRO 770 48.913 43.329 65.392 1.00 1.00 0 ATOM 5470 CPRO 770 48.913 43.329 65.392 1.00 1.00 0 ATOM 5471 CD PRO 770 46.081 47.016 64.640 1.00 2.00 0 ATOM 5472 N ASN 771 48.363 47.259 64.138 1.00 2.00 0 ATOM 5478 CD ASN 771 48.00 49.664 65.022 1.00 40.66 0 ATOM 5478 CD ASN 771 48.00 49.664 65.022 1.00 40.66 0 ATOM 5478 DD ASN 771 48.00 49.664 65.022 1.00 40.66 0 ATOM 5478 DD ASN 771 50.811 49.318 65.208 1.00 50.00 0 ATOM 5478 DD ASN 771 50.811 49.318 65.208 1.00 50.00 0 ATOM 5478 DD ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5480 CD ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5481 C ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5482 C ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5483 N TYR 772 50.657 47.191 66.417 1.00 6.11 0 ATOM 5481 C ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5480 CD TYR 772 51.555 46.645 64.477 1.00 6.11 0 ATOM 5481 C ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5482 C ASN 771 50.814 48.414 64.666 1.00 2.00 0 ATOM 5483 N TYR 772 50.657 47.191 66.466 1.00 2.00 0 ATOM 5480 CD TYR 773 55.634 64.647 66.81 1.00 6.11 0 ATOM 5480 CD TYR 7							38.645	65.091	1.00 2.00	0
ATOM 5458 O SER 768 46.297 40.731 65.336 1.00 2.00 0 ATOM 5458 O SER 768 46.297 40.731 65.973 1.00 2.00 0 ATOM 5459 N ALA 769 44.615 41.911 65.973 1.00 2.00 0 ATOM 5461 CA ALA 769 44.615 41.911 65.973 1.00 2.00 0 ATOM 5462 CB ALA 769 44.619 43.009 66.428 1.00 27.69 0 ATOM 5463 C ALA 769 45.142 42.454 67.786 1.00 4.86 0 ATOM 5463 C ALA 769 45.140 44.253 66.018 1.00 31.25 0 ATOM 5465 N PRO 770 45.717 44.109 65.786 1.00 11.84 0 ATOM 5466 CD PRO 770 46.717 44.109 65.786 1.00 11.84 0 ATOM 5466 CD PRO 770 47.556 29.914 65.393 1.00 2.00 0 ATOM 5468 CB PRO 770 46.717 44.109 65.786 1.00 14.07 0 ATOM 5468 CB PRO 770 46.717 41.09 65.786 1.00 14.07 0 ATOM 5469 CG PRO 770 46.717 41.09 65.786 1.00 14.07 0 ATOM 5467 CA ARM 771 46.101 64.401 0.00 0.00 0 ATOM 5471 C PRO 770 46.221 46.784 65.393 1.00 2.00 0 ATOM 5472 N ASM 771 46.813 46.303 65.393 1.00 1.00 2.00 0 ATOM 5473 N ASM 771 48.703 65.394 1.00 1.00 2.00 0 ATOM 5474 C A ASM 771 48.703 46.612 46.400 0.00 0.00 0 ATOM 5475 CB ASM 771 48.703 47.556 64.550 1.00 2.00 0 ATOM 5475 CB ASM 771 48.703 47.556 64.550 1.00 2.00 0 ATOM 5476 CB ASM 771 48.703 47.556 64.550 1.00 2.00 0 ATOM 5478 ND2 SSN 771 48.270 51.032 64.457 1.00 2.00 0 ATOM 5478 CB ASM 771 48.203 51.0357 64.550 1.00 40.36 0 ATOM 5478 CB ASM 771 48.203 51.032 64.457 1.00 47.17 0 ATOM 5488 CB ASM 771 49.320 51.032 64.457 1.00 47.17 0 ATOM 5488 CB ASM 771 49.320 51.032 64.457 1.00 47.17 0 ATOM 5488 CB ASM 771 50.174 48.414 64.666 1.00 2.00 0 ATOM 5488 CB ASM 771 50.174 48.414 64.666 1.00 2.00 0 ATOM 5488 CB ASM 771 50.811 49.318 65.208 1.00 50.90 0 ATOM 5488 CB ASM 771 50.811 49.318 65.208 1.00 50.90 0 ATOM 5488 CB ASM 771 50.811 49.318 65.208 1.00 60.00						43.903	37.635			0
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ATOM 5528 CZ PHE 776 56.845 46.833 60.071 1.00 6.06 0										
								58.969		
ATUM 5529 C PHE 7/6 51.424 50.402 60.460 1.00 21.71 0										
	ATOM	5529	Ċ	PHE	116	51.424	JU.4UZ	50.480	1.00 21.71	U

MOTA	5530	0	PHE	776	50.979	50.601	59.346	1.00 9.78	0
MOTA	5531	N	ASP	777	50.757 49.427	50.720 51.322	61.584 61.532	1.00 47.34 1.00 47.73	0
MOTA	5533	CA	ASP ASP	7 7 7 7 7 7	49.526	52.790	61.071	1.00 36.82	ŏ
ATOM ATOM	5534 5535	CB CG	ASP	777	50.299	53.677	62.059	1.00 86.03	0
ATOM	5536	OD1		777	51.519	53.464	62.246	1.00 85.84	0
ATOM	5537	OD2	ASP	777	49.685	54.595	62.644	1.00 85.92 1.00 47.70	0
ATOM	553B	C	ASP	7 7 7	48.456 47.458	50.547 51.095	60.632 60.1 64	1.00 47 .70 1.00 37.69	0
MOTA	5539	0	ASP	7 7 7 7 7 8	48.756	49.268	60.419	1.00 12.97	ŏ
MOTA MOTA	5540 5542	N CA	asn asn	778	47.948	48.373	59.584	1.00 3.97	Ŏ
ATOM	5543	CB	ASN	778	48.760	47.123	59.221	1.00 17.34	0
ATOM	5544	CG	ASN	778	49.081	46.258	60.435 61.275	1.00 11.19 1.00 10.18	0
MOTA	5545	OD1		778	49.928 48.404	46.613 45.120	60.538	1.00 7.69	ŏ
MOTA	5546 5549	ND2 C	asn asn	778 778	46.672	47.945	60.296	1.00 4.32	Ō
MOTA MOTA	5550	ŏ	ASN	778	46.497	48.227	61.481	1.00 21.92	0
ATOM	5551	N	ALA	779	45.783	47.267	59.578	1.00 6.26 1.00 6.26	0
MOTA	5553	CA	ALA	779	44.537 43.404	46.799 46.923	60.168 59.174	1.00 6.26 1.00 78.54	Ö
MOTA	5554	СВ	ALA	779 779	44.727	45.344	60.580	1.00 6.26	ŏ
ATOM ATOM	5555 5556	C O	ALA ALA	779	45.537	44.626	59.988	1.00 86.19	0
ATOM	5557	N	GLY	780	43.998	44.925	61.606	1.00 2.00	0
MOTA	5559	CA	GLY	780	44.091	43.560	62.077 61.747	1.00 2.00 1.00 2.00	0
MOTA	5560	C	GLY	780 780	42.800 41.725	42.848 43.433	61.869	1.00 21.77	ŏ
MOTA	5561 5562	O N	GLY ALA	781	42.880	41.587	61.345	1.00 2.00	0
MOTA MOTA	5564	CA	ALA	781	41.670	40.870	60.984	1.00 2.00	0
ATOM	5565	CB	ALA	781	41.671	40.567 39.587	59.492 61.763	1.00 14.58 1.00 2.00	0
ATOM	5566	C	ALA	781 781	41.474 42.396	39.076	62.398	1.00 14.58	Õ
ATOM	5567 5568	O N	ALA MET	782	40.263	39.061	61.660	1.00 2.00	0
MOTA MOTA	5570	CA	MET	782	39.857	37.833	62.319	1.00 2.00	0
ATOM	5571	CB	MET	782	39.037	38.204	63.554 64.502	1.00 2.00 1.00 2.00	0
MOTA	5572	CG	MET	782	38. 7 36 37. 74 3	37.067 37.608	65.912	1.00 2.00	ŏ
ATOM	5573	SD CE	MET MET	782 782	37.595	39.370	65.621	1.00 2.00	0
MOTA MOTA	5574 5575	C	MET	782	39.010	37.029	61.306	1.00 2.00	0
ATOM	5576	ō	MET	782	38.090	37.570 35.752	60. 68 8 61.116	1.00 2.00 1.00 2.00	ő
MOTA	5577	N	MET	783	39. 32 5 38. 57 0	34.955	60.165	1.00 2.00	Õ
MOTA	5579 5580	CA CB	MET	783 783	39,482	34.182	59.209	1.00 16.43	0
MOTA MOTA	5581	CG	MET	783	38. 68 8	33.363	58.165	1.00 19.34 1.00 18.69	0
MOTA	5582	SD	MET	783	39.689	32.337 30.793	57.060 57.9 7 5	1.00 21.53	ŏ
MOTA	5583	CE	MET	783 783	39.780 37.608	33.976	60.794	1.00 2.00	0
MOTA MOTA	5584 5585	C O	met Met	783	38.004	33.010	61.447	1.00 10.33	0
MOTA	5586	N	SER	784	36.335	34.221	60.5 4 2 61.034	1.00 2.00 1.00 2.00	0
MOTA	5588	CA	SER	784	35.276 34.033	33.374 34.220	61.308	1.00 16.25	ō
MOTA	5589	CB	SER	784 784	34.385	35.392	62.020	1.00 16.25	0
MOTA MOTA	5590 5592	OG C	SER SER	784	34.953	32.301	59.990	1.00 2.00	0
ATOM	5593	ŏ	SER	784	34.672	32.611	58.839 60.401	1.00 19.28 1.00 10.88	Ö
MOTA	5594	N	VAL	785	35.033 34.706	31.045 29.907	59.557	1.00 16.53	0
MOTA	5596	CA	VAL	785 785	35.649	28.739	59.792	1.00 11.43	0
MOTA	5597 5598	CB CG1	VAL VAL	785	35.183	27.538	58.975	1.00 11 43	0
MOTA MOTA	5599	CG2		785	37.082	29.153	59.477	1.00 11.43	0
ATOM	5600	С	VAL	785	33.357	29.465 28.998	60.083 61.225	1.00 12.30	Ö
MOTA	5601	0	VAL	785 786	33.254 32.307	29.613	59.291	1.00 2.00	0
MOTA	5602	N CA	ASP ASP	786 786	31.024	29.200	59.807	1.00 2.00	0
MOTA MOTA	5604 5605	CB	ASP	786	29.874	30.008	59.171	1.00 24.90 1.00 30.71	0
ATOM	5606	CG	ASP	786	29.403	29.458 29.765	57.842 57.474	1.00 33.80	Ö
MOTA	5607	OD:		786 786	28.245 30.165	28.737	57.163	1.00 32.56	0
ATOM	5608	OD:	2 ASP ASP	786	30.860	27.699	59.656	1.00 2.00	0
	EEAA								
MOTA MOTA	5609 5610	C	ASP	786	31.677	27.025	59.031	1.00 23.30	0

					20 003	27.195	60.268	1.00 7.48	٥
MOTA	5611	N	GLU	787	29.803 29.447	25.782	60.270	1.00 13.41	Ö
MOTA	5613	CA	GLU	787 787	27.983	25.636	60.696	1.00 2.00	0
MOTA	5614	CB	GLU	787	27.174	26.963	60.747	1.00 2.00	0
MOTA	5615	CG	GLU GLU	787	27.384	27.758	62.053	1.00 2.00	0
MOTA	5616	CD	GTU	787	26.858	27.319	63.117	1.00 2.00	0
MOTA	5617	OE2	GLU	787	28.070	28.817	62.024	1.00 2.00	0
MOTA	5618 5619	C	GLU	787	29.665	25.015	58.965	1.00 11.86	0
MOTA MOTA	5620	ŏ	GLU	787	30.002	23.830	58.994	1.00 2.00	0
ATOM	5621	Ň	THR	788	29.492	25.698	57.836	1.00 56.74	0
ATOM	5623	CA	THR	788	29.616	25.092	56.513	1.00 52.70	0
ATOM	5624	CB	THR	788	28.369	25.402	55.700	1.00 6.46	0
ATOM	5625	OG1	THR	788	28.317	26.819	55.467	1.00 4.65	0
MOTA	5627	CG2	THR	78 8	27.101	24.968	56.457	1.00 10.48 1.00 51.86	0
MOTA	5628	С	THR	788	30.835	25.536	55.683	1.00 12.46	0
MOTA	5629	0	THR	788	30.751	25.628	54.449 56.364	1.00 10.40	ő
ATOM	5630	N	LEU	789	31.946	25.823 26.245	55.734	1.00 6.34	ŏ
MOTA	5632	CA	LEU	789	33.194	25.165	54.775	1.00 8.80	ŏ
MOTA	5633	CB	LEU	789	33.670 34.458	24.017	55.405	1.00 15.93	ŏ
MOTA	5634	CG	LEU	789	35.879	24.487	55.677	1.00 11.55	ō
MOTA	5635		LEU	789 789	33.785	23.525	56.682	1.00 14.98	Ō
MOTA	5636		LEU	789	33.173	27.619	55.042	1.00 4.53	0
MOTA	5637	C	LEU	789	34.065	27.948	54.248	1.00 8.97	0
MOTA	5638	O N	MET	790	32.165	28.430	55.336	1.00 2.00	0
MOTA	5639 5641	CA	MET	790	32.126	29.748	54.743	1.00 2.00	0
MOTA MOTA	5642	CB	MET	790	30.698	30.267	54.636	1.00 19.79	0
ATOM	5643	CG	MET	790	30.588	31.572	53.882	1.00 19.37	0
ATOM	5644	SD	MET	790	28.979	31.708	53.134	1.00 18.51	0
ATOM	5645	CE	MET	790	28.359	33.110	53.992	1.00 23.23	0
ATOM	5646	C	MET	790	32. 94 5	30.667	55.627	1.00 2.00	0
ATOM	5647	0	MET	790	32.615	30.857	56.799	1.00 23.79 1.00 2.00	0
MOTA	5648	N	CYS	791	34.014	31.228	55.067	1.00 2.00 1.00 2.00	o
MOTA	5650	CA	CYS	791	34.882	32.133	55.803 55.516	1.00 14.83	ŏ
MOTA	5651	CB	CYS	791	36.325	31.793	55.766	1.00 15.43	õ
ATOM	5652	SG	CYS	791	36.570	30.052 33.591	55.502	1.00 2.00	ō
MOTA	5653	Ċ	CYS	791	34.616 34.314	33.961	54.371	1.00 21.84	Ō
MOTA	5654	0	CYS	791 792	34.697	34.412	56.540	1.00 2.00	0
MOTA	5655	N	SER	792	34.480		56.430	1.00 2.00	0
MOTA	5657	CA	SER SER	792	33.073		56.903	1.00 2.93	0
MOTA	5658 5659	CB OG	SER	792	32.887		58.256	1.00 3.41	0
MOTA MOTA	5661	C	SER	792	35.539		57.303	1.00 2.00	0
ATOM	5662	Ö	SER	792	36.290	35.826	57.994	1.00 2.56	0
ATOM	5663	Ň	PHE	7 9 3	35.609	37.842	57.270	1.00 2.00	0
ATOM	5665	CA	PHE	793	36.617		58.039	1.00 2.00	0
ATOM	5666	CB	PHE	793	37.765		57.129	1.00 38.05 1.00 24.74	ő
ATOM	5667	CG	PHE	793	38.531		56.469	1.00 24.74 1.00 28.08	ő
MOTA	5668		PHE	793	38.055		55.306	1.00 26.11	ŏ
MOTA	5669		PHE	793	39.7 4 1 38.774		57.002 5 4.6 80	1.00 27.84	Ō
MOTA	5670		PHE	793	40.466		56.389	1.00 27.16	0
MOTA	5671	CE2		793 793	39.983		55.225	1.00 29.41	0
ATOM	5672	CZ	PHE	793	36.101		58.784	1.00 2.00	0
MOTA	5673 5674	CO	PHE	793	35.362		58.233	1.00 23.22	0
MOTA	5675	N	GLN	794	36.480		60.051	1.00 67.38	0
MOTA MOTA	5677	CA	GLN	794	36.128		60.837	1.00 63.52	0
MOTA	5678	CB	GLN	794	35.608		62.23B	1.00 31.74	0
MOTA	5679	CG	GLN	794	34.294	39.954	62.289	1.00 36.03	0
MOTA	5680	CD	GLN	794	34.506		62.252	1.00 39.99	0
MOTA	5681	OE		794	34.622		61.179	1.00 39.14	0
MOTA	5682	NE		794	34.575	37.838	63.425	1.00 44.72	0
MOTA	5685	C	GLN	794	37.471		60.958	1.00 65.96	0
MOTA	5686	ō	GLN	794	38.486		61.272	1.00 37.24 1.00 53.07	0
MOTA	5687	N	ILE	795	37. 49 8		60.658	1.00 54.05	0
MOTA	5689	CA	ILE	795	38.732		60.748 59.392	1.00 26.19	0
MOTA	5690	CB	ILE	795	39.084		59.608	1.00 28.70	0
MOTA	5691	CC:	2 ILE	795	40.115	45.589	,,.000	1.00 20.70	J

ATOM MOTA MOTA MOTA MOTA MOTA MOTA MOTA	5692 5693 5694 5695 5696 5698 5700 5701 5702 5703 5704 5705 5707 5709 5710 5711	CG1 CD1 C O N CA CB CGCD1 CD2 C O N CA CB CCD CE		795 795 795 796 796 796 796 796 796 797 797 797	39.643 38.659 38.543 37.476 39.544 39.539 40.976 39.242 40.637 41.649 40.455 40.805 39.456 39.134	43.467 42.449 44.847 45.452 45.008 46.054 45.528 44.121 44.038 43.114 46.977 46.510 48.280 49.207 50.354 49.929 51.183 50.875	58.390 57.906 61.817 61.903 62.674 63.686 65.089 65.308 66.702 65.119 63.206 62.667 63.356 62.902 62.154 60.932 60.151 58.662 57.851	1.00 30.47 1.00 28.83 1.00 53.94 1.00 27.08 1.00 34.91 1.00 41.88 1.00 41.67 1.00 32.51 1.00 33.78 1.00 34.59 1.00 37.84 1.00 40.12 1.00 0.26 1.00 0.34 1.00 0.34	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5712 5716 5717 5718 5720 5721 5722 5723 5724 5726 5727 5728 5729 5730 5732	NZ C O N CA C C O N CA C C C C C C C C C C C C C C C C C C	LYS LYS ALA ALA ALA ALA ARG ARG ARG ARG ARG ARG	797 797 797 400 400 400 401 401 401 401 401 401	38.852 42.356 42.961 -8.399 -6.981 -6.530 -7.350 -5.218 -4.619 -4.586 -5.638 -5.638 -6.285 -6.338	33.565 33.160 34.922 35.762 35.116 36.351 36.395 37.360 37.340 38.512 39.260 40.369	64.053 63.985 131.469 131.818 130.576 132.352 132.744 132.833 134.359 134.956 136.471 137.058 138.022 138.457	1.00 50.42 1.00 0.89 1.00 75.56 1.00 75.56 1.00 21.27 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00	000000000000000000000000000000000000000
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5733 5736 5739 5741 5741 5744 5745 5746 5749 5751 5755 5755	NH12 CONCACBCG1 CG2 CONCACBCG2 CONCACBCG	ARG ARG VAL VAL VAL VAL	401 401 402 402 402 402 402 402 403 403 403 403	-4.527 -3.216 -2.505 -2.822 -1.516 -1.331 -0.063 -0.398 -0.548 0.701 1.882 1.894 1.503 3.070 2.890	38.949 36.446 35.443 37.655 37.882 39.379 39.632 39.889 37.376 36.920 36.461 34.942 34.298	138.511 132.267 132.150 131.898 131.311 131.042 130.283 130.282 132.223 133.444 131.625 132.361 131.333 3 131.514	1.00 2.00 1.00 2.50 1.00 23.92 1.00 23.92 1.00 23.72	00000000000000000
MOTA MOTA MOTA MOTA MOTA MOTA MOTA MOTA	5757 5759 5760 5761 5762 5763 5764 5766 5766 5769 5771 5772 5773	N CA CB CC	PHE PHE PHE PHE PHE PHE PHE PHE ALA ALA ALA ALA	404 404 404 404 404 404 404 405 405 405	4.271 5.439 6.101 5.228 4.421 5.171 3.560 4.313 3.507 6.476 6.472 7.341 8.420 9.762 8.434 8.380 32.968	36.827 38.060 39.270 39.574 40.086 41.17 41.47 35.73 34.72 35.95 35.96 33.68 32.65 37.22	4 132.711 0 130.505 8 132.677 8 130.463 2 131.549 5 130.791 0 129.802 129.414 2 129.619 2 130.078 4 129.401 6 49.661	1.00 13.47 1.00 13.06 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 2.00 1.00 85.03 1.00 95.94 1.00 95.94	000000000000000000000000000000000000000

	585 5	^	OT Y	000	31.215	18.274	51.038	1.00 95.94	0
MOTA	5778	Č	GLY	8 9 9 89 9	30.022	18.368	51.338	1.00 32.75	ŏ
MOTA	5779	0	GLY ARG	900	32.090	19.260	51.193	1.00 42.23	ŏ
MOTA	5780	N CA	ARG	900	31.732	20.569	51.722	1.00 42.23	Ō
MOTA	5782 5783	CB	ARG	900	31.110	20.432	53.110	1.00 22.01	0
MOTA MOTA	5784	ÇG	ARG	900	30.578	21.718	53.694	1.00 22.01	0
ATOM	5785	CD	ARG	900	30.080	21.488	55.120	1.00 22.01	0
ATOM	5786	NE	ARG	900	31.005	20.664	55.909	1.00 22.01	0
ATOM	5788	CZ	ARG	900	31.111	20.691	57.237	1.00 22.01	0
ATOM	5789	NH1		900	30.361	21.512	57.963	1.00 22.01	0
ATOM	5792		ARG	900	31.957	19.865	57.839	1.00 22.01	0
ATOM	5795	С	ARG	900	33.034	21.365	51.774	1.00 42.23	0
MOTA	5796	0	ARG	900	33.795	21.309	52.735	1.00 22.01	0
MOTA	5797	N	ARG	901	33.284	22.091	50.697	1.00 8.39	0
MOTA	5799	CA	ARG	901	34.490	22.880	50.542	1.00 8.39	0
MOTA	5800	CB	ARG	901	34.793	23.036	49.045	1.00 2.00	0
ATOM	5801	CG	ARG	901	34.504	21.810	48.186	1.00 2.00	0
ATOM	5802	CD	ARG	901	34.584	22.175	46.699	1.00 2.00	0
MOTA	5803	NE	ARG	901	34.278	21.012	45.863	1.00 2.00 1.00 2.00	0
MOTA	5805	CZ	ARG	901	35.190	20.195	45.327	1.00 2.00 1.00 2.00	0
ATOM	5806		ARG	901	36.498	20.416	45.509 44.649	1.00 2.00	Ö
MOTA	5809	NH2	ARG	901	34.791 34.327	19.114 24.259	51.158	1.00 2.00	Ö
MOTA	5812	C	ARG	901	33.241	24.629	51.607	1.00 2.00	ő
ATOM	5813	0	ARG	901	35.393	25.005	51.149	1.00 2.00	ő
MOTA	5814	N	VAL	902	35.425	26.384	51.650	1.00 2.00	ŏ
MOTA	5816	CA	VAL	902 902	36.880	26.814	51.869	1.00 12.90	Õ
ATOM	5817	CB CG1	VAL	902	37.039	28.332	51.992	1.00 12.90	ŏ
MOTA	5 81 8	CG2	VAL	902	37.492	26.219	53.139	1.00 12.90	Ö
MOTA	5819 5820	CGZ	VAL	902	34.782	27.324	50.627	1.00 2.00	Ō
ATOM ATOM	5821	Ö	VAL	902	34.737	27.031	49.435	1.00 12.90	0
MOTA	5822	N	SER	903	34.288	28.438	51.120	1.00 2.00	0
ATOM	5824	CA	SER	903	33.673	29.487	50.271	1.00 2.00	0
ATOM	5825	CB	SER	903	32.173	29.205	50.008	1.00 2.00	0
ATOM	5826	0G	SER	903	31.477	28.917	51.209	1.00 2.00	0
ATOM	5828	c	SER	903	33.871	30.814	50.991	1.00 2.00	0
MOTA	5829	ō	SER	903	34.386	30.866	52.115	1.00 2.00	0
ATOM	5830	N	PHE	904	33.482	31.903	50.376	1.00 2.00	0
MOTA	5832	CA	PHE	904	33.683	33.188	51.034	1.00 2.00	0
ATOM	5833	CB	PHE	904	34.894	33.898	50.455	1.00 17.14	0
MOTA	5834	CG	PHE	904	36.184	33.109	50.672	1.00 17.14 1.00 17.14	0
MOTA	5835		PHE	904	36.511	32.067	49.805	1.00 17.14 1.00 17.14	ő
ATOM	5836	CD2	PHE	904	37.034	33.426	51.738 49.994	1.00 17.14	Ö
ATOM	5837		PHE	904	37.690 38.216	31.343 32.704	51.927	1.00 17.14	Õ
MOTA	5838	CE2	PHE	904	38.544	31.662	51.054		Ö
ATOM	5839	CZ	PHE	904	32.491	34.080	50.894	1.00 2.00	ō
MOTA	5840	C	PHE PHE	904 904	31.716	33.994	49.926	1.00 17.14	0
MOTA	5841 5842	O N	ALA	905	32.386	34.912	51.874	1.00 70.83	0
MOTA	5844	CA	ALA	905		35.857	51.949	1.00 69.99	0
ATOM ATOM	5845	CB	ALA	905	30.042	35.128	52.370	1.00 2.00	0
ATOM	5846	C	ALA	905	31.646	36.940	52.962	1.00 69.85	0
ATOM	5847	Õ	ALA	905	30.981	37.068	53.993	1.00 2.00	0
ATOM	5848	N	ALA	907	37.374	34.312	47.285	1.00 23.39	0
ATOM	5850	CA	ALA	907	36.215	33.651	46.695	1.00 23.39	0
MOTA	5851	CB	ALA	907	35.317	34.681	46.008	1.00 41.64	0
MOTA	5852	Ċ	ALA	907	36.637	32.568	45.699	1.00 23.39	0
ATOM	5853	0	ALA	907	35. 85 0	31.680	45.370	1.00 41.64	0
ATOM	5854	N	ALA	908	37.875	32.643	45.215	1.00 83.90	0
MOTA	5856	CA	ALA	908	38.378	31.663	44.253	1.00 83.90	0
ATOM	5857	CB	ALA	908	39.439	32.296	43.337	1.00 22.84	0
ATOM	5858	С	ALA	90B	3B.961	30.466	44.992	1.00 83.90	0
MOTA	5859	0	ALA	908	38.459	30.074	46.046	1.00 22.84	0
MOTA	5860	OW	TAW	1	62.869	37.982	63.341 61.228	1.00 20.00 1.00 20.00	0
MOTA	5863	OW	WAT	103	57.039 7.257	39.062 66.194	118.365	1.00 20.00	Ö
MOTA	5866	OW	WAT	101	12.341		123.464	1.00 20.00	0
MOTA	5869	OW	WAT	2 3	10.655		120.833	1.00 20.00	õ
MOTA	5872	OW	WAT	,	.0.0.	30.740		30.00	-

ATOM	5875	OW	WAT	104	55.432	36.306	63.901	1.00 20.00	0
ATOM	5878	OW	WAT	102	56. 78 2	40.457	58.333	1.00 20.00	0
	5881	OW	WAT	4	5. 78 7	57.856	118.686	1.00 20.00	0
MOTA	5884	OW	WAT	105	54.382	39.155	63.734	1.00 20.00	0
ATOM			WAT	5	8.964	57.595	118.151	1.00 20.00	0
MOTA	5887	OW			38.565	47.423	74.959	1.00 20.00	ŏ
MOTA	5890	OW	WAT	106	16.086	42.169	105.289	1.00 20.00	ŏ
MOTA	5893	OM	TAW	6				1.00 20.00	Ö
ATOM	5896	OW	WAT	107	31.158	26.414	51.913		
ATOM	5899	OW	WAT	7	-0.781	32.787	131.574	1.00 20.00	0
MOTA	5902	MN	MN2	430	4.422	59.061	119.360	1.00 15.61	0
ATOM	5903	MN	MN2	431	7.458	57.875	117.661	1.00 16.53	0
	5904	MIN	MN2	930	56.038	34.500	63.727	1.00 16.67	0
MOTA		MN	MN2	931	54.402	37.798	64.756	1.00 15.40	0
MOTA	5905		S04	801	57.551	37.278	64.009	1.00 37.87	0
MOTA	5906	S.		801	57.600	35.852	63.897	1.00 42.46	Ó
MOTA	5907	01	S04		58.690	37.740	64.722	1.00 42.01	Ŏ
ATOM	5908	02	SO4	801			64.705	1.00 45.08	ŏ
MOTA	5909	03	S04	801	56.355	37.648			ő
MOTA	5910	04	S04	801	57.520	37.854	62.725	1.00 41.40	_
ATOM	5911	S	SO4	800	6.866	60.776	118.643	1.00 37.87	0
ATOM	5912	01	SO4	800	7.710	60.635	119.773	1.00 42.46	0
		02	SO4	800	7.044	62.063	118.053	1.00 42.01	0
MOTA	5913			800	5.496	60.612	119.046	1.00 45.08	0
MOTA	5914	03	S04		7.194	59.728	117.703	1.00 41.40	0
MOTA	5915	04	S04	800	1.134	37.120			-

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CLAIMS

- 1. A method of identifying a compound which modulates the interaction between a PP1c and a regulatory subunit thereof, the method comprising determining whether a compound enhances or disrupts the interaction between (a) a PP1c or a fragment, variant, derivative or fusion thereof or a fusion of a fragment, variant or derivative and (b) a regulatory subunit which is able to bind to PP1c or a PP1c-binding fragment, variant, derivative or fusion of said subunit or a fusion of said fragment, variant or derivative.
 - 2. A method of identifying a compound which mimics the effect of a regulatory subunit of PP1c, the method comprising contacting said compound with PP1c and determining whether, in the presence of the compound, PP1c adopts the function of properties of a PP1c in the presence of a given regulatory subunit.
- 3. A method according to Claim 1 or 2 wherein said regulatory subunit of PP1c is any one of M₁₁₀, G_L, G_M, M-complexes, p53 BP2, sds22, NIPPI, L5, Inhibitor-1, Inhibitor-2, or DARPP.
 - 4. A method according to Claim 3 wherein the regulatory subunit of PP1c is any one of M_{110} , G_L , G_M , M-complexes or p53BP2.
- 25 5. A method according to Claim 4 wherein the regulatory subunit of PP1c is M_{110} or G_M .
- 6. A method according to Claim 1 wherein the fragment of a regulatory subunit which is able to bind to PP1c is any of the peptides [G63-T93], [G63-N75], [E2-P243], [E2-D118], and peptide 63-80 of G_M or

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functional equivalents thereof or peptides comprising said peptide sequences provided that they are not the complete G_M regulatory subunit.

- 7. A method according to Claim 1 wherein the fragment of a regulatory subunit which is able to bind to PP1c is any of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M₁₁₀ or functional equivalents thereof or peptides comprising said peptide sequences provided that they are not the complete M₁₁₀ regulatory subunit.
- A method according to Claim 1 wherein the PP1c-binding fragment, variant or derivative of said regulatory subunit or a fusion of said fragment, variant or derivative comprises the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any amino acid.
- 15 9. A method according to Claim 8 wherein the PP1c-binding fragment, variant or derivative comprises, in addition to the said consensus peptide sequence, at least one basic residue in the four residues N-terminal of the consensus peptide sequence.
- 20 10. A method according to Claim 8 wherein in the consensus peptide sequence Xaa is not Asp or Glu or a large hydrophobic residue.
 - 11. A method according to Claim 8 wherein the PP1c-binding fragment is a fragment of a regulatory subunit comprising the said consensus peptide sequence.
 - A method according to Claim 10 wherein the PP1c-binding fragment is a fragment of any of the M₁₁₀, G_L, G_M, M-complexes, p53BP2, sds22, NIPPI, L5, Inhibitor-1, Inhibitor-2 or DARPP regulatory subunits comprising said consensus sequence.

- 13. A method according to any one of the preceding claims wherein the compound binds to a PP1c.
- 14. A method according to Claim 1 wherein the compound binds to a regulatory subunit of PP1c.
 - 15. A compound identifiable by the method of any one of Claims 1 to 14.
- 16. A compound which modulates the interaction between a PP1c and a regulatory subunit thereof said compound comprising any of the peptides [G63-T93], [G63-N75], [E2-P243], [E2-D118], and peptide 63 to 80 of G_M or functional equivalents or said compound comprising any of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M₁₁₀ or functional equivalents thereof or said compound comprising the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any naturally occurring amino acid or functional equivalents thereof, provided that the said compound is not a complete regulatory subunit of PP1c.
- 17. A compound according to Claim 16 consisting of the peptides [G63-T93], [G63-N75], [E2-P243], [E2-D118], or peptide 63 to 80 of G_M or functional equivalents thereof or consisting of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M₁₁₀ or functional equivalents thereof.
 - 18. A peptide capable of binding a PP1c and which affects the ability of PP1c to bind to a particular target and/or affects the regulation of PP1c activity, or a functional equivalent thereof.
- 30 19. A method of identifying a compound which modulates the interaction

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between a PP1c and a regulatory subunit thereof, or binds PP1c or mimics the effect of a regulatory subunit, the method comprising selecting a compound which is capable of adopting the same or substantially the same conformation as a peptide bound to the regulatory subunit-binding site of PP1c or the same or substantially the same conformation as the portion of PP1c which binds to said peptide.

- 20. A method according to Claim 19 wherein said peptide comprises the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe wherein Xaa is any amino acid.
 - 21. A method according to Claim 20 wherein said peptide consists of residues 63 to 75 of G_M .
- A method according to Claim 21 wherein the conformation of the said peptide and the conformation of the said portion of PP1c is as defined by reference to the coordinates in Table A.
 - 23. A compound identifiable by the method of any one of Claims 19 to 22.
 - 24. A compound according to any one of Claims 15 to 18 or 23 for use in medicine.
- 25. A pharmaceutical composition comprising a compound according to any one of Claims 15 to 18 or 23 and a pharmaceutically acceptable carrier.
 - 26. A method of affecting cellular metabolism or function, the method comprising administering to a cell (a) a compound which modulates the interaction between a PP1c and a regulatory subunit thereof or (b) a compound which mimics the effect of a regulatory subunit of PP1c or

- (c) a peptide capable of binding a PP1c and which affects the ability of PP1c to bind to a particular target and/or affects the regulation of PP1c activity, or a functional equivalent thereof.
- A method according to Claim 26 wherein any one or more of the peptides [G63-T93], [G63-N75], [E2-P243], [E2-D118], and peptide 63-80 of G_M or functional equivalents thereof or peptides comprising said peptide sequences are administered.
- 10 28. A method according to Claim 26 wherein any one or more of the peptides [M1-E309], [M1-F38], [M1-A150] or [L24-Y496] of M₁₁₀ or functional equivalents thereof or peptides comprising said peptide sequences are administered.
- 15 29. A method according to Claim 26 wherein a compound according to any one of Claims 15 to 18 or 23 are administered to the cell.
 - 30. A method according to any one of Claims 26 to 29 wherein the cell is in a mammalian body.
 - 31. A method of treating a patient in need of modulation of PP1c activity or function the method comprising administering to the patient an effective amount of a compound which modulates the interaction between a PP1c and a regulatory subunit thereof or (b) a compound which mimics the effect of a regulatory subunit of PP1c or (c) a peptide capable of binding a PP1c and which affects the ability of PP1c to bind to a particular target and/or affects the regulation of PP1c activity, or a functional equivalent thereof.
- 30 32. Use of peptides derived from targeting subunits of PP1c, functional

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equivalents or portions thereof to affect cellular metabolism.

- 33. A method of treatment of a mammal said method comprising altering levels of peptides derived from a targeting subunit of PP1c, functional equivalents or portions thereof to an extent that cellular metabolism or function is affected.
- 34. A PP1c-regulating subunit that is modified so that it cannot interact with PP1c.
- 35. A PP1c-regulator subunit according to Claim 34 wherein the consensus peptide sequence Arg/Lys-Val/Ile-Xaa-Phe is missing or modified.

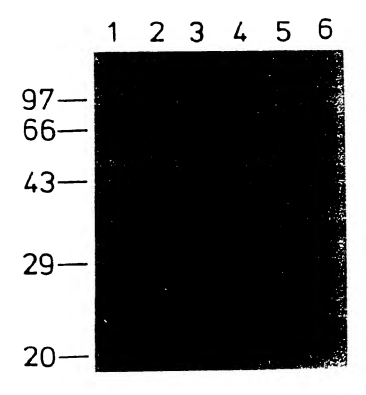


Fig. 1

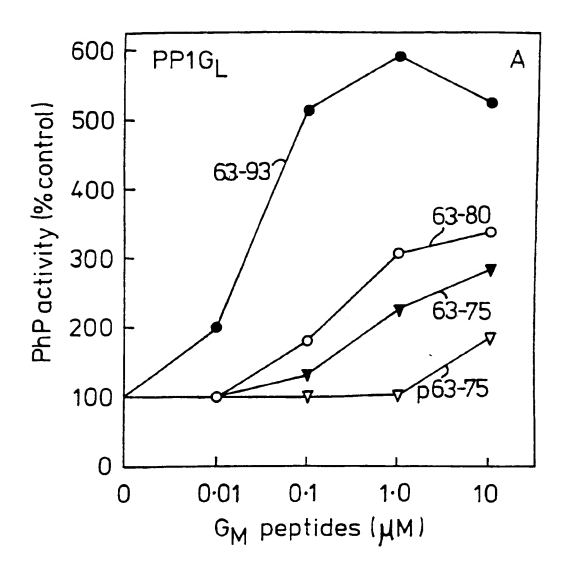


Fig. 2

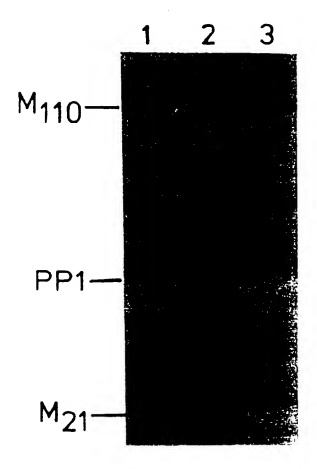
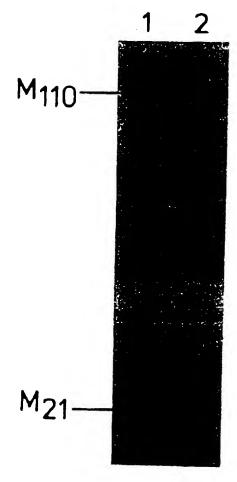


Fig. 3A



MP:PhP 0-92 0-97

Fig. 3B

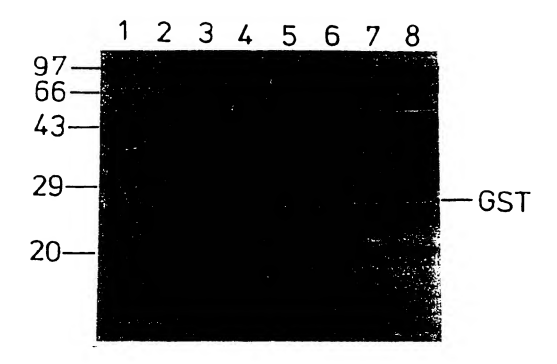


Fig. 4

0.001 0.01

0.1

1

M₁₁₀ fragment (nM)

10

100

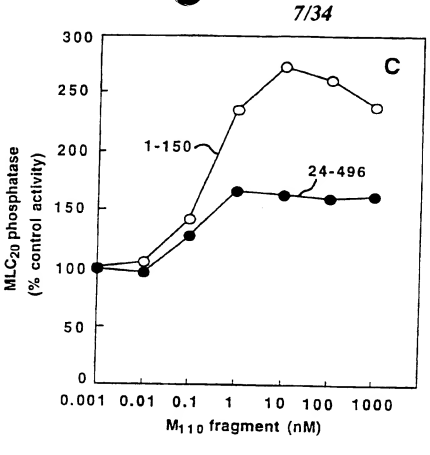


Fig. 5C

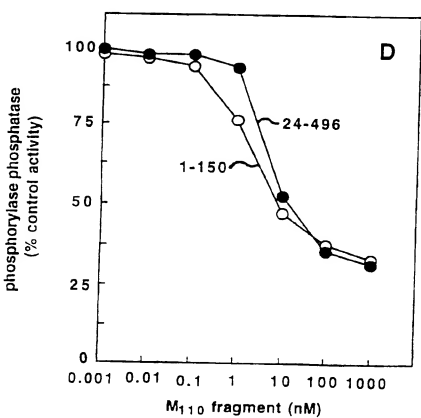


Fig. 5D

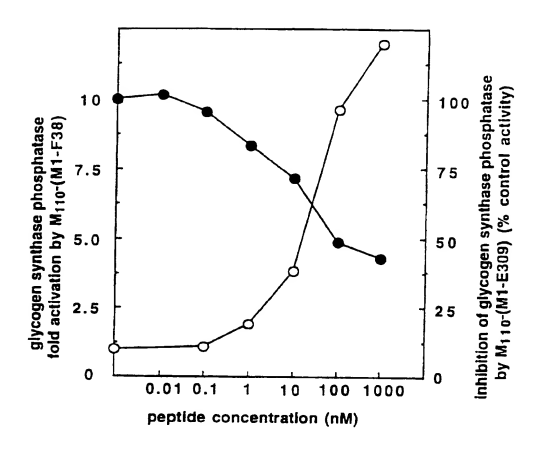


Fig. 6

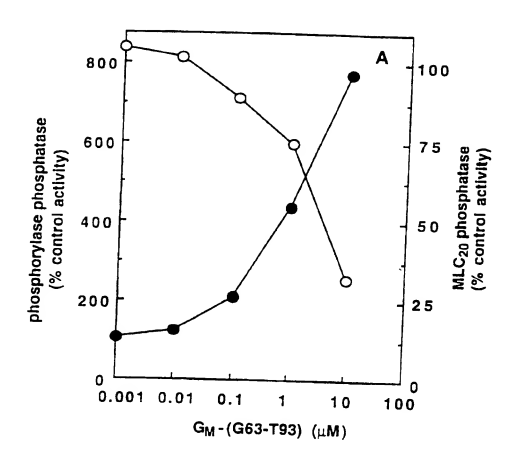


Fig. 7A

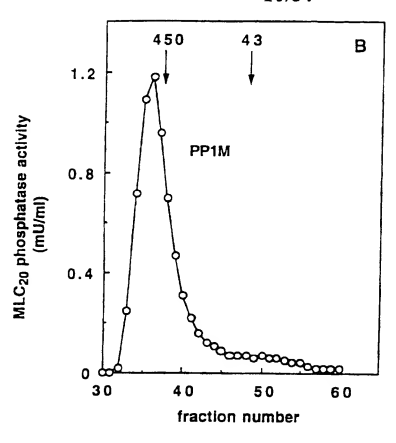


Fig. 7B

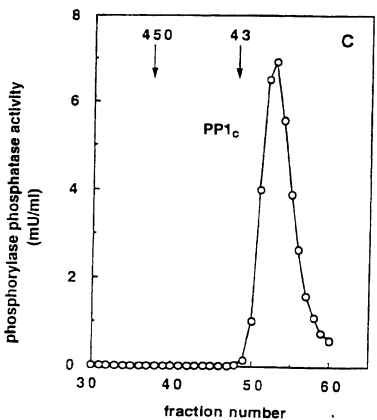
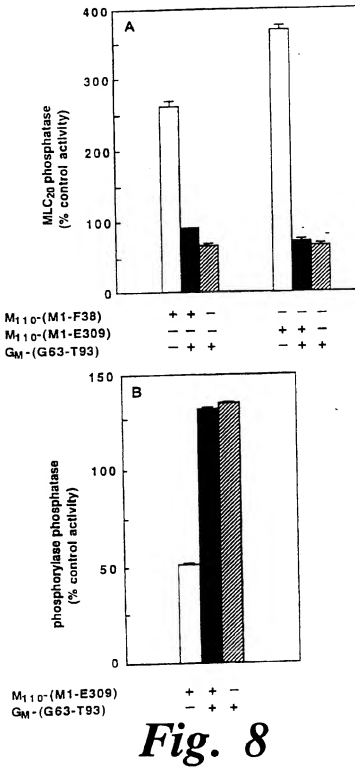


Fig. 7*C*



Effect of fragment on $PP1G_L$ and/or PP1M activity

and/or PP1M by fragment Dissociation of-PPIG_L

prevents G_L from suppressing the

G63 🗆 T93

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G63[] N75

does not dissociate G_L from PP1G_L displaces M₁₁₀ from PP1M

dephosphorylation of phosphorylase dephosphorylation of phosphorylase prevents G_L from suppressing the

displaces G_L, from PP1G_L

MICHARININE ESHIP

MI BESSE A 150

M1 | F38

MIL NEWS BEST

M₁₁₀

dephosphorylation of phosphorylase dephosphorylation of phosphorylase MLC20 but does not suppress the stimulates dephosphorylation of stimulates dephosphorylation of stimulates dephosphorylation of MLC20 and suppresses MLC₂₀ and suppresses

no effect on PP1C activity

D39 KKKK KKKU F.309

dephosphorylation of phosphorylase

displaces G_L from PP1G_L



Fig. 10A

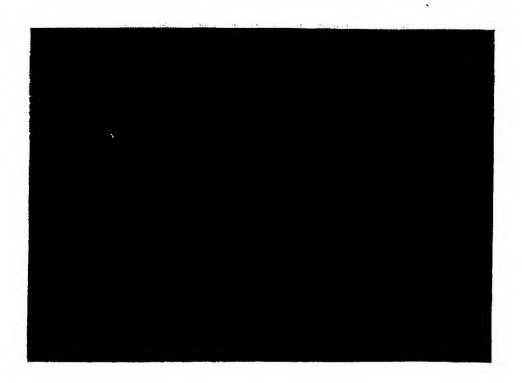


Fig. 10B

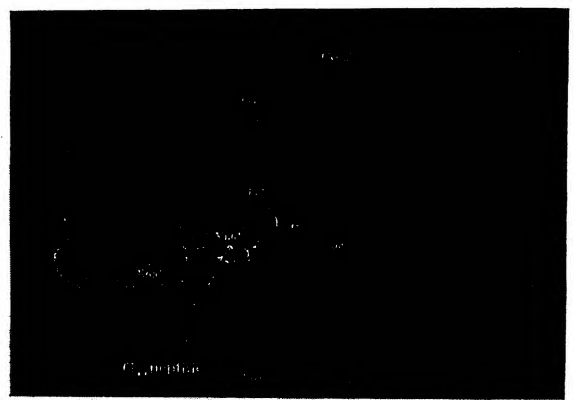


Fig. 11A

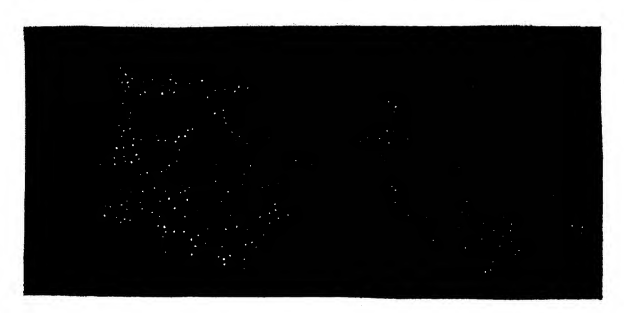


Fig. 11B

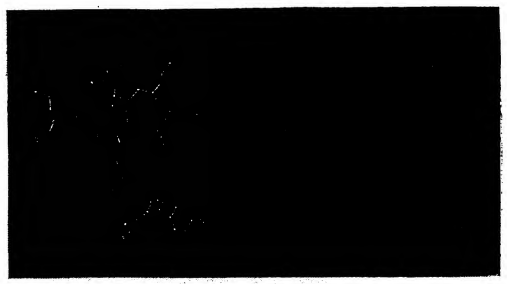


Fig. 11C

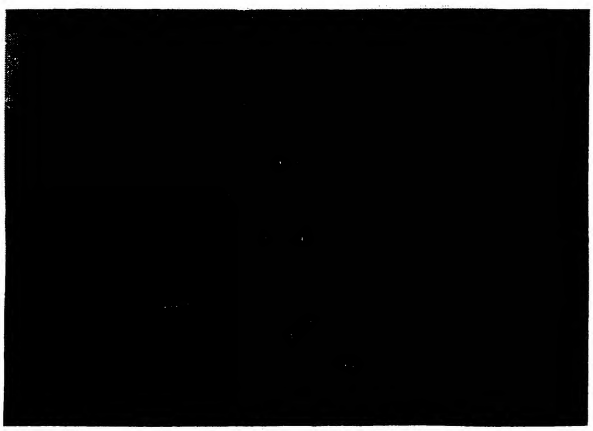


Fig. 11E

protein	putative PP1-binding motif	residues
	,•	
GAC1	SPEKNVRFAIE	66-76
PIG2	S S G K S V R F A A H	50-60
GIP2	IRSKSVHFDQA	217-227
YIL045W	Q R S K S V H F D R V	193-203
YIL045W	V F V K N I Y F S N A	412-422
REG1	TKNRHIHFNDR	461-471
REG2	PRERHIK F NDN	164-174
SCD5	FKSKKVRFSEH	270-280
GIP1	LSEKFIPFNNL	180-190
GIP1	KKK RCVNF RNK	441-451
SHP1	KVTREITFWKE	232-242

Fig. 12A

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protein	Puc	at	iv	e	PP	1-1	bi	nd	in	g 1	not	if residues
GAC1	s	P	Ε	K	N	v	R	F	A	Ī	Ε	66-76
PIG2	S	s	G	ĸ	s	v	R	F	A	A	H	50-60
GIP2	I	R	s	ĸ	s	v	H	F	D	Q	A	217-227
YIL045W	Q	R	s	ĸ	s	v	H	F	D	R	v	193-203
YIL045W	V	F	v	ĸ	N	I	Y	F	s	N	A	412-422
REG1	Ŧ	K	N	R	Ħ	I	Ξ	F	N	D	R	. 461-471
REG2	7	₽.	Ξ	₹	Ħ	I	ĸ	F	N	ם	N	164-174
SCD5	F	K	s	ĸ	ĸ	v	R	F	s	Ξ	H	270-280
GIP1	W	N	L :	X	F	I	P	F	И	N	L	180-190
GIP1	K	K :	K [1	R (<u> </u>	<u>۷</u> 1	N :	F	R	N	ĸ	441-451

Fig. 12B

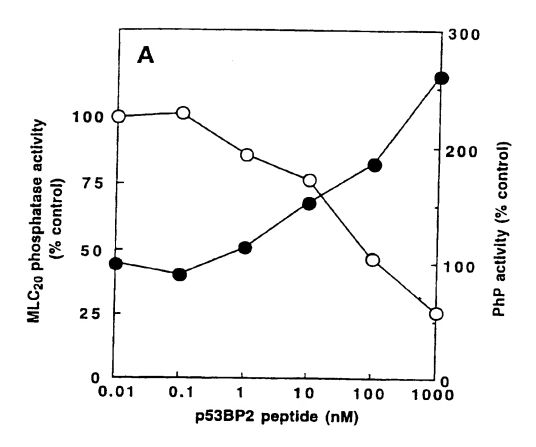


Fig. 13A

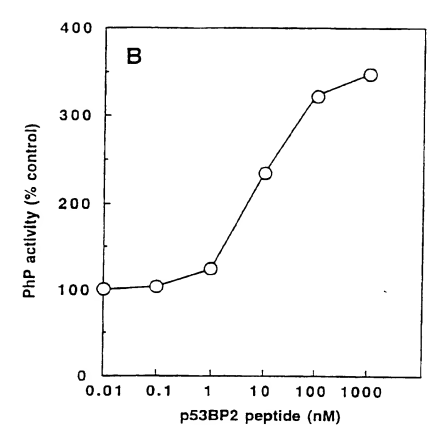


Fig. 13B

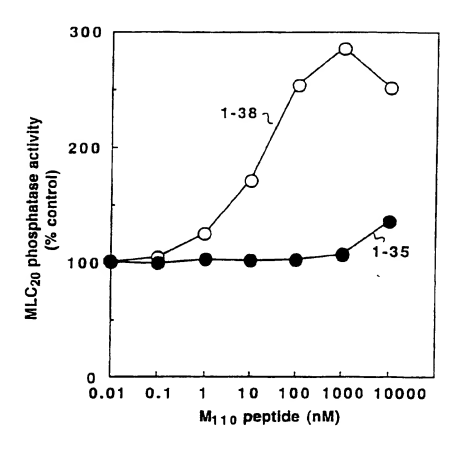


Fig. 14

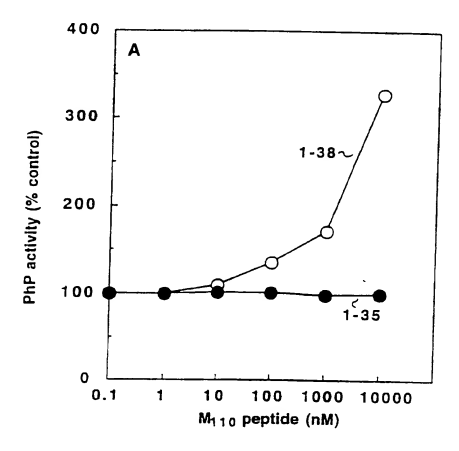


Fig. 15A

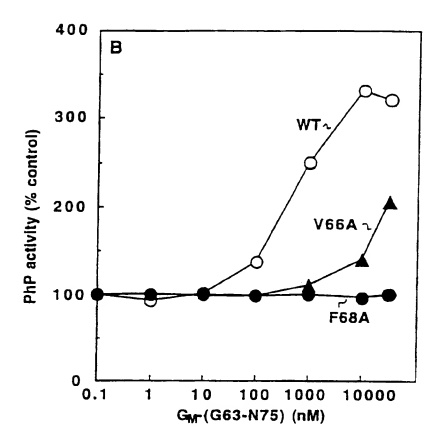


Fig. 15B

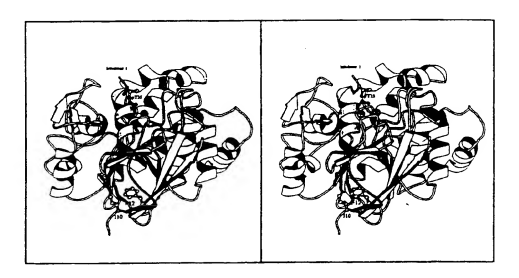


Fig. 16

Rat	HEMADAKQKRWEQLERMIGBETDLEPPVYKRQKTEVKPDDGAVFLAACSS		Fig.	17
Rat Ch	GDTDEVLKLLERGADI MYAMVDGLTALEQACI DDMVDMVKFLVEMGAMIM : : :		1 15.	 /
Rat	OPDWIGWIPLEARSCGYLDIAEFLIGOGREY GRYMSEGDTPLDIAEEER	150		
Ch Rat	Opdnegwipleaaasc gyldiaeylisogaev gavnsegdtpldiaeer Heellonevnrogydieaarkeeerihlrdar onlusgeisdvreaksgg			
СÞ	HEELLQHEVERQGVDI EAARKEEERIMLRDAR QWLHSGEIMDVREAKSGG			
Ch Ch	TALEVARARGYTEVLKLLIQAGYDVMIKDYDG WYPLEARAEWGKEERCRI 	250 250		
Rat Ch	Lydnlcdmetverygqtafdyadedilgtlerlorkghlesekrdkesp : Lyenlcdmeaymrygqtafdyadedilgtlerlorkghlesekrrkksp	300 300		
Rat	LIESTAMMENNOPORTPRHEETLIIRPERNAS RIESLEGEKADEELEGKK ::		*	
Rat	DESCSSEEDEEDDSESEARTDETERMASVIRAETASTQAAPAAVTTPTL			
Ch Rat	DESSCHEEEDDDSESSAETDRAKTLA NAMTTSTQSASMTAPSV SENGGTPTSPVKKFPTSTTKISPKEEERKDES PASWRLGLEKTGSTGALA	395 450		
Ch	AGGGGTPTSPLEXFPTSTTEVSPEEREREDES PASWELGLERETGSTGALA	445		
Rat	ZITASKEAQKEKDTAGVIRSASSPRLSSSLDE KEKERDHKGTRLAYVAPT	500 495		
Rat	IPRREGSTSDIEEKEMRESSMLRTSSSYTR REWEDDLEEMSSIMEGST	548		
Ch Rat	ipriletediderem <u>rdssassirsgasyar remezdykka. Slargp</u> t	544 552/593	1	
CJ7	YERSCSYGRRODDLISCSYPSTTSTPT VISAAGLOKSPLESTSTT	592		
RAE Ch	AKTPPGSSPAGTOSST SMRLWARDSTEKERDS APTARTILVAPTVVSAAA	587/643 642		
Rat	SSTTALTTTAGTLESTSEVRERREYLTPVR DEESESQREARSROAROS	637/693		
Ch.	. TITANTTATSQTVSS TSEVRERRRSTLTPVR DEESEEQREARSRQARQS RRSTQGVTLTDLQEARXTIGRSRSTRTREQEN ENDREEREKQDKEKQFE	691 687/743		
Ch	RESTOGOTION OF A STRUCTURE RESTRICT RES	741		
Rat	KERSEVSREDEYKÇKYSETYDETYARYRPVST SSSSTPSSSSLSTLGSSL	737/793		
Rat	KKKSE. TKDDDYRGRISKTVEEPTERIRPTST. STSTSSTSSLSTSTSSL YASSQLERPESLVOITSAYSROLTKDWEREGE KKEEEKEGEDKSQPKSIR	789 787/843		
C).	### PART OF THE PA	*31		
Rat Ch	EXRAPREERRSIGYSFWTQDSDENEQERQSDT EDGSSKRDTQTDSVSETD	837/893		
Rat	SSSTSSDRYDSLLGRSASYSYLEERKPYGSRLEEDDSTDFKKLYZQI			
Ch Ratl/3	tgáláváágdátbaggátággátládriþyc átláredátdþitlitgi Erra	938		
Rat2 Ch	LARMEKLEAQLEDTHMEETDLELQLERATORO ERFADESLLEMEKETOR 	935		
Rat1/3 Rat2	LERRIBENEEELKMLPDLKADNQRLKDEWGAL IRVISELEK SQYLLGGTKSSRKKNI	976/103 951	2	
CÞ	SQYLLGGKRSSRKEDI	1004		

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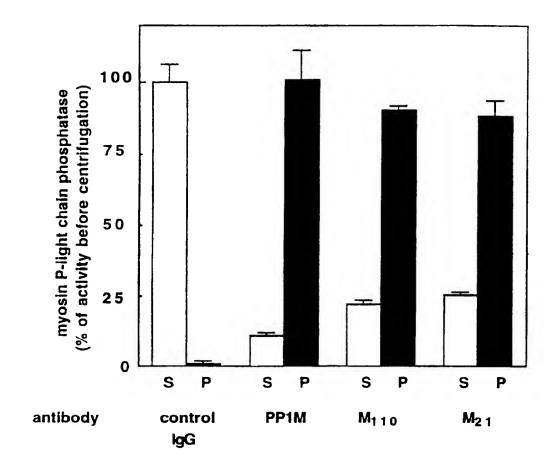


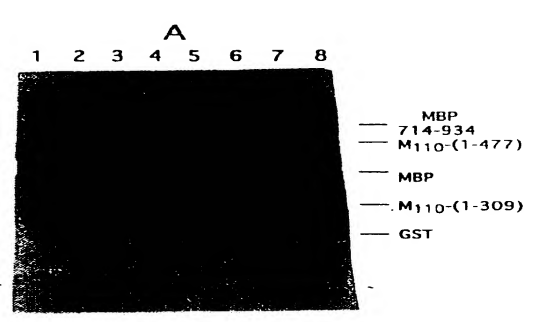
Fig. 18A

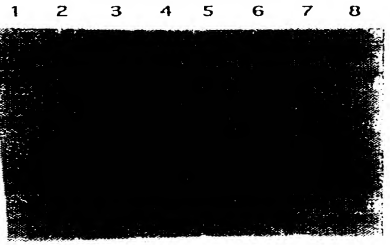






Fig. 18B





В

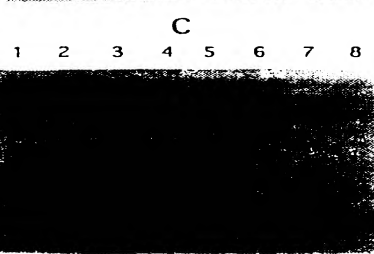


Fig. 19

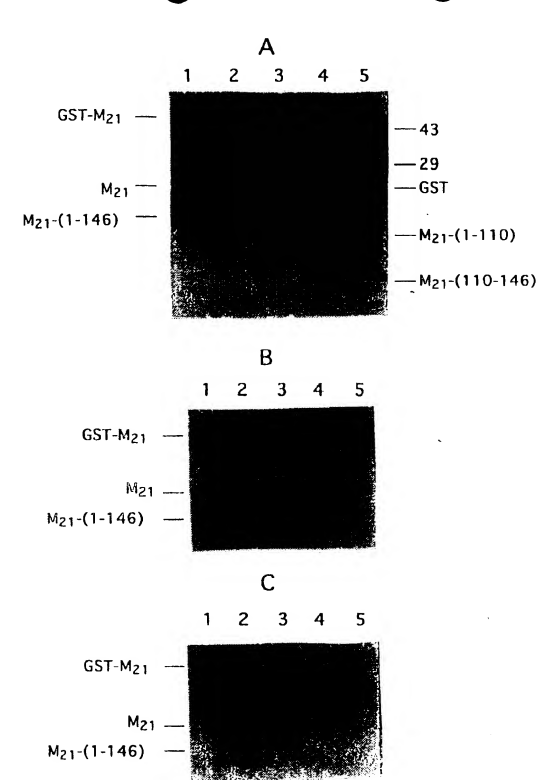


Fig. 20

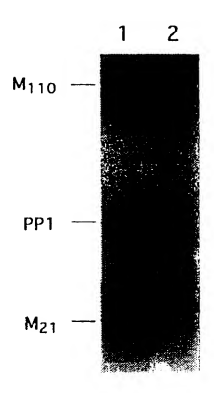


Fig. 21

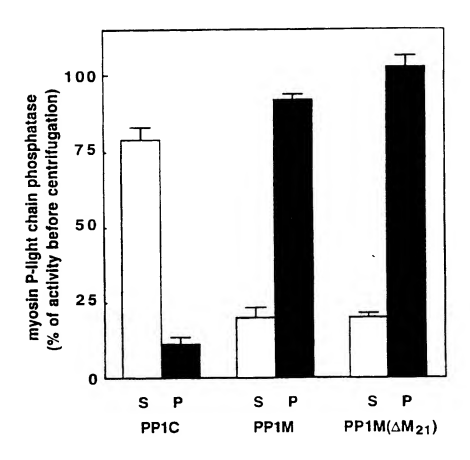
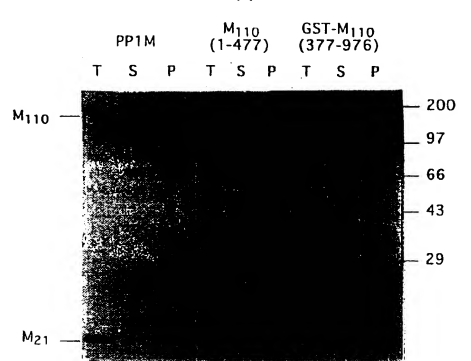


Fig. 22

A



B

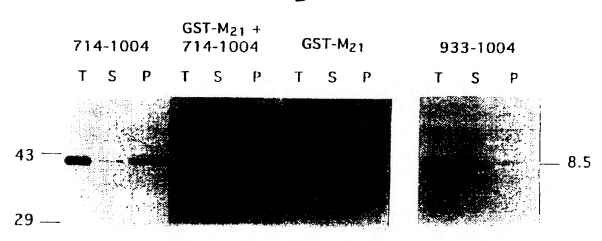


Fig. 23

A

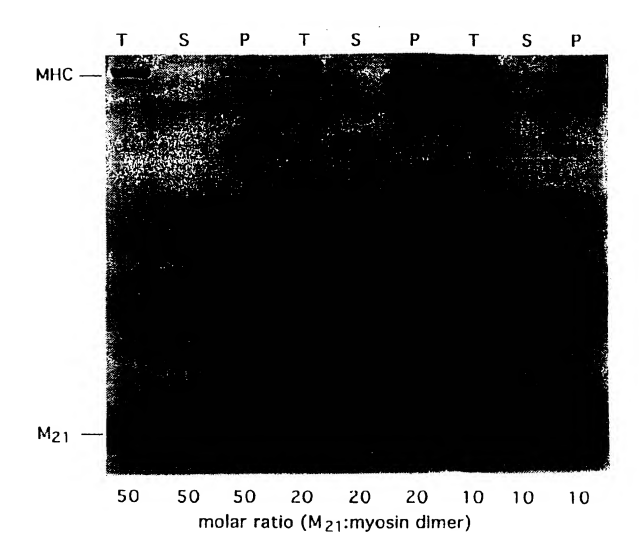


Fig. 24A

В

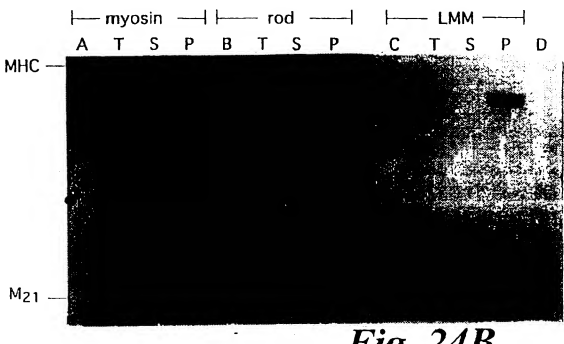


Fig. 24B С

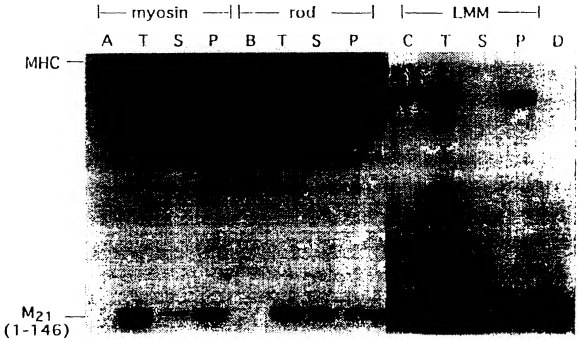


Fig. 24C

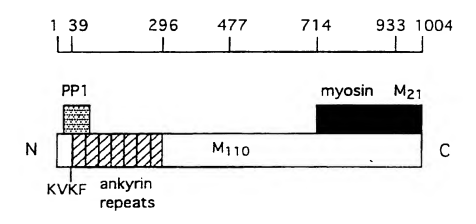


Fig. 25

INTERNATIONAL SEARCH REPORT

intbonal	Application No
GB	97/00898

			7			
A. CLASSIFICATION OF SUBJECT MATTER IPC 6 G01N33/573 C12N9/12 C07K14/435						
According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED						
Minimum d IPC 6	locumentation as withed (classification system followed by classification GOIN CIZN CO7K	tion symbols)				
Documental	tion searched other than minimum documentation to the extent that	such documents are included in the fields s	earched			
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)						
	MENTS CONSIDERED TO BE RELEVANT					
Calegory *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claum No.			
A	TRENDS IN BIOCHEMICAL SCIENCE, vol. 18, 1 May 1993, OXFORD UK, pages 172-177, XP002037474 M.J. HUBBARD ET AL.: "On target new mechanism for the regulation protein phosphorylation" cited in the application see the whole document JOURNAL OF MOLECULAR BIOLOGY,		1-35			
	vol. 254, 1995, NEW YORK NY USA, pages 942-959, XP002037475 M.P. EGLOFF ET AL.: "Crystal st the catalytic subinit of human p phosphatase 1 and its complex wi tungstate." see the whole document	rotein				
X Furt	her documents are listed in the continuation of box C.	Patent family members are listed in	n annex.			
	* Special categories of cited documents: "T' later document published after the international filing date or priority date and not in conflict with the application but cited to independ on the priorities or theory importance the					
considered to be of particular relevance invention						
filing of "L" docume which citation	date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another n or other special reason (as specified)	"X" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is taken alone. "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the				
'O' document referring to an oral disclosure, use, exhibition or other means document is combined with one or more other such document, such combination being obvious to a person skilled in the art. A' document member of the same patent family						
	Date of the actual completion of the international search Date of mailing of the international search					
13 August 1997						
Name and n	nailing address of the ISA European Patent Office, P.B. 5818 Patentiann 2	Authorized officer				
	NL - 2220 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Van Bohemen, C				

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INTERNATIONAL SEARCH REPORT

Int attonal Application No P 97/00898

		3 97/00898
	abon) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	CRITICAL REVIEWS IN BIOCHEMISTRY AND MOLECULAR BIOLOGY, vol. 27, no. 3, 1992, BOCA RATON FL USA, pages 227-281, XP002037649 M. BOLLEN ET AL.: "The structure and role and regulation of type 1 protein phosphatase." cited in the application see the whole document	1-35
T	EMBO JOURNAL, vol. 16, no. 8, 1997, HEIDELBERG FRG, pages 1876-1887, XP002037247 M.P. EGLOFF ET AL.: "Structural basis for the recognition of regulatory subunits by the catalytic subunit of protein phosphatase 1." see page 1876, column 2, line 10 - line 49	1-35

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